

10/660,118

=&gt; d his nofile

(FILE 'HOME' ENTERED AT 14:47:35 ON 23 FEB 2006)

FILE 'CAPLUS' ENTERED AT 14:47:57 ON 23 FEB 2006

 SET LINE 250  
 SET DETAIL OFF  
 E US2003-660118/AP,PRN 25  
 SET NOTICE 1000 SEARCH  
 L1 1 SEA ABB=ON US2003-660118/AP  
 SET NOTICE LOGIN SEARCH  
 SET LINE LOGIN  
 SET DETAIL LOGIN  
 D SCAN  
 SEL RN

FILE 'REGISTRY' ENTERED AT 14:50:02 ON 23 FEB 2006

 L2 21 SEA ABB=ON (117525-18-5/BI OR 117525-19-6/BI OR 3483-12-3/BI  
 OR 53-57-6/BI OR 616-91-1/BI OR 675214-32-1/BI OR 675214-33-2/B  
 I OR 675214-34-3/BI OR 675214-35-4/BI OR 675214-36-5/BI OR  
 675214-37-6/BI OR 675214-38-7/BI OR 675214-39-8/BI OR 675214-40  
 -1/BI OR 675214-41-2/BI OR 675214-42-3/BI OR 675214-43-4/BI OR  
 675625-84-0/BI OR 675625-85-1/BI OR 70-18-8/BI OR 9074-14-0/BI)

D SCAN

FILE 'CAPLUS' ENTERED AT 14:53:26 ON 23 FEB 2006

 L3 2951 SEA ABB=ON WHITE C?/AU  
 L4 4538 SEA ABB=ON THIOREDOXIN#/OBI  
 L5 9 SEA ABB=ON L3 AND L4  
 L6 2350 SEA ABB=ON SPUTUM/CT  
 L7 4370 SEA ABB=ON MUCUS/CT  
 L8 9174 SEA ABB=ON CYSTIC FIBROSIS/OBI  
 L9 27 SEA ABB=ON L4 AND (L6 OR L7 OR L8)  
 L10 9 SEA ABB=ON L4 AND (L6 OR L7)  
 D SCAN  
 L11 3325 SEA ABB=ON THIOREDOXINS/CT  
 L12 7 SEA ABB=ON L11 AND (L6 OR L7)  
 L13 728128 SEA ABB=ON 9/SC, SX  
 L14 4 SEA ABB=ON L12 NOT L13  
 L15 3 SEA ABB=ON L12 AND L13  
 D SCAN  
 L16 472 SEA ABB=ON MUCOLY?/OBI  
 L17 29200 SEA ABB=ON LIQUEF?/OBI  
 E LIQUIDIFI/CT  
 E LIQUIDIFI/BI  
 L18 161373 SEA ABB=ON VISCO?/OBI  
 L19 1055 SEA ABB=ON EXPECTORANT#/OBI  
 L20 12 SEA ABB=ON L11 AND L8 NOT L13  
 L21 8 SEA ABB=ON L20 NOT (L1 OR L5 OR L14)  
 D SCAN  
 L22 7029 SEA ABB=ON CYSTIC FIBROSIS/CT  
 L23 5 SEA ABB=ON L22 AND L11 NOT (L13 OR L1 OR L5 OR L14)  
 D SCAN TI

FILE 'REGISTRY' ENTERED AT 15:19:04 ON 23 FEB 2006

L24 541003 SEA ABB=ON .C..C./SQSP

FILE 'CAPLUS' ENTERED AT 15:19:39 ON 23 FEB 2006

FILE 'REGISTRY' ENTERED AT 15:19:48 ON 23 FEB 2006  
D RN L24 270000  
L25 271004 SEA RAN=(,518362-12-4) ABB=ON .C..C./SQSP  
L26 269999 SEA ABB=ON L24 NOT L25

FILE 'CAPLUS' ENTERED AT 15:21:37 ON 23 FEB 2006  
L27 58456 SEA ABB=ON L25 OR L26  
L28 389 SEA ABB=ON L27 AND L11  
L29 82 SEA ABB=ON L27 AND (L6 OR L7)  
L30 8 SEA ABB=ON L29 AND (L16 OR L17 OR L18 OR L19)  
D SCAN TI

FILE 'REGISTRY' ENTERED AT 15:25:54 ON 23 FEB 2006  
L31 5728 SEA ABB=ON CGPC/SQSP

FILE 'CAPLUS' ENTERED AT 15:26:20 ON 23 FEB 2006  
L32 2177 SEA ABB=ON L31  
L33 8 SEA ABB=ON L32 AND (L6 OR L7)  
L34 2 SEA ABB=ON L30 AND L33  
D SCAN TI L33

FILE 'MEDLINE' ENTERED AT 15:27:56 ON 23 FEB 2006  
E THIOREDOXIN/CT  
E E3+ALL  
L35 2163 SEA ABB=ON THIOREDOXIN/CT  
L36 2181 SEA ABB=ON WHITE C?/AU  
L37 7 SEA ABB=ON L35 AND L36  
D TRIAL 1-7  
L38 11646 SEA ABB=ON SPUTUM/CT  
L39 13832 SEA ABB=ON VISCOSITY/CT  
E MUCUS+ALL/CT  
L40 8697 SEA ABB=ON MUCUS+NT/CT  
L41 1 SEA ABB=ON L35 AND L39 AND (L38 OR L40)  
L42 3 SEA ABB=ON L35 AND (L38 OR L39 OR L40)  
D TRIAL 1-3  
L43 19620 SEA ABB=ON CYSTIC FIBROSIS/CT  
L44 2 SEA ABB=ON L43 AND L35

FILE 'EMBASE' ENTERED AT 15:31:48 ON 23 FEB 2006  
L45 1505 SEA ABB=ON WHITE C?/AU  
E THIOREDOXIN+ALL/CT  
L46 2331 SEA ABB=ON THIOREDOXIN/CT  
E MUCUS+ALL/CT  
L47 6283 SEA ABB=ON MUCUS+NT/CT  
E SPUTUM+ALL/CT  
L48 3562 SEA ABB=ON SPUTUM/CT  
E VISCOSITY+ALL/CT  
L49 100 SEA ABB=ON SPUTUM VISCOSITY/CT  
L50 3436 SEA ABB=ON VISCOELASTICITY/CT  
L51 9749 SEA ABB=ON VISCOSITY/CT  
L\*\*\* DEL 8 S L45 AND L46  
D TRIAL 1-8  
L52 267 SEA ABB=ON MUCOLYSIS/CT  
L53 183 SEA ABB=ON LIQUEFACTION/CT  
L54 8 SEA ABB=ON L45 AND L46  
L55 5 SEA ABB=ON L46 AND (L47 OR L48 OR L49 OR L50 OR L51 OR L52 OR L53)  
E CYSTIC FIBROSIS+ALL/CT  
L56 20389 SEA ABB=ON CYSTIC FIBROSIS/CT  
L57 7 SEA ABB=ON L46 AND L56

L58 4 SEA ABB=ON L57 NOT (L54 OR L55)  
D TRIAL 1-4  
L59 1 SEA ABB=ON L46(L) DT/CT AND L56

FILE 'DRUGU' ENTERED AT 15:36:48 ON 23 FEB 2006  
L60 356 SEA ABB=ON WHITE C?/AU  
E THIOREDOXIN/CT  
L61 72 SEA ABB=ON THIOREDOXIN#/CT  
L62 2 SEA ABB=ON L60 AND L61  
D TRIAL 1-2  
L63 405 SEA ABB=ON CYSTIC-FIBROSIS/CT  
L64 1059 SEA ABB=ON SPUTUM/CT  
L65 4493 SEA ABB=ON MUCOLYTIC#/CT  
E MUCUS/CT  
L66 972 SEA ABB=ON MUCUS/CT  
E VISCOSITY/CT  
E E3+ALL  
L67 2111 SEA ABB=ON VISCOSITY/CT  
E LIQUEF/CT  
L68 14 SEA ABB=ON LIQUEFACTION/CT OR LIQUEFYING/CT  
L69 3 SEA ABB=ON L61 AND (L63 OR L64 OR L65 OR L66 OR L67 OR L68)  
D TRIAL 1-3  
L70 45091 SEA ABB=ON RESPIRATORY/CC  
L71 1 SEA ABB=ON L61 AND (L63 OR L64 OR L65 OR L66 OR L67 OR L68)  
AND L70

FILE 'STNGUIDE' ENTERED AT 15:40:45 ON 23 FEB 2006

FILE 'JICST-EPLUS, PASCAL, WPIX, IPA, BIOSIS, ESBIOBASE, BIOTECHDS,  
LIFESCI, CONFSCI, DISSABS, SCISEARCH' ENTERED AT 15:47:40 ON 23 FEB 2006

L72 11489 SEA ABB=ON WHITE C?/AU  
L73 17233 SEA ABB=ON THIOREDOXIN#  
L74 38537 SEA ABB=ON SPUTUM  
L75 47183 SEA ABB=ON MUCUS  
L76 3898 SEA ABB=ON MUCOLY?  
L77 74427 SEA ABB=ON LIQUEF?  
L78 567528 SEA ABB=ON VISCO?  
L79 79051 SEA ABB=ON CYSTIC FIBROSIS  
L80 10 SEA ABB=ON L72 AND L73 AND (L74 OR L75 OR L76 OR L77 OR L78  
OR L79)  
L81 8 SEA ABB=ON L73 AND L76  
L82 14 SEA ABB=ON L73 AND (L74 OR L75) AND (L77 OR L78 OR L79)

FILE 'STNGUIDE' ENTERED AT 15:50:22 ON 23 FEB 2006

FILE 'CAPLUS' ENTERED AT 15:51:08 ON 23 FEB 2006

D QUE L1  
D QUE L5

L83 9 SEA ABB=ON L1 OR L5

FILE 'MEDLINE' ENTERED AT 15:51:08 ON 23 FEB 2006

D QUE L37

FILE 'EMBASE' ENTERED AT 15:51:08 ON 23 FEB 2006

D QUE L54

FILE 'DRUGU' ENTERED AT 15:51:08 ON 23 FEB 2006

D QUE L62

FILE 'JICST-EPLUS, PASCAL, WPIX, IPA, BIOSIS, ESBIOBASE, BIOTECHDS,

LIFESCI, CONFSCI, DISSABS, SCISEARCH' ENTERED AT 15:51:25 ON 23 FEB 2006  
D QUE L80

FILE 'MEDLINE, DRUGU, CAPLUS, EMBASE, PASCAL, WPIX, BIOSIS, ESBIOBASE,  
BIOTECHDS, SCISEARCH' ENTERED AT 15:51:39 ON 23 FEB 2006

L84 20 DUP REM L37 L62 L83 L54 L80 (16 DUPLICATES REMOVED)  
ANSWERS '1-7' FROM FILE MEDLINE  
ANSWERS '8-9' FROM FILE DRUGU  
ANSWERS '10-13' FROM FILE CAPLUS  
ANSWERS '14-16' FROM FILE EMBASE  
ANSWERS '17-18' FROM FILE PASCAL  
ANSWERS '19-20' FROM FILE ESBIOBASE  
D IALL 1-9  
D IBIB ED ABS HITIND 10-13  
D IALL 14-20

FILE 'MEDLINE' ENTERED AT 15:52:20 ON 23 FEB 2006

FILE 'STNGUIDE' ENTERED AT 15:52:47 ON 23 FEB 2006

FILE 'REGISTRY' ENTERED AT 15:53:28 ON 23 FEB 2006  
D QUE L24

FILE 'CAPLUS' ENTERED AT 15:53:28 ON 23 FEB 2006  
D QUE L30

L85 6 SEA ABB=ON L30 NOT L83  
D IBIB ED ABS HITRN L85 1-6  
SEL HIT RN L85 1-6

FILE 'STNGUIDE' ENTERED AT 15:54:51 ON 23 FEB 2006

FILE 'REGISTRY' ENTERED AT 15:55:07 ON 23 FEB 2006

L86 98 SEA ABB=ON (132053-08-8/BI OR 143831-71-4/BI OR 132053-07-7/BI  
OR 182177-07-7/BI OR 182177-08-8/BI OR 182177-09-9/BI OR  
182177-10-2/BI OR 182177-11-3/BI OR 182177-12-4/BI OR 182177-13  
-5/BI OR 182177-14-6/BI OR 182177-15-7/BI OR 182177-16-8/BI OR  
182177-17-9/BI OR 182177-18-0/BI OR 182177-19-1/BI OR 182177-20  
-4/BI OR 182177-21-5/BI OR 182177-22-6/BI OR 182177-23-7/BI OR  
182177-24-8/BI OR 182177-25-9/BI OR 182177-26-0/BI OR 182177-27  
-1/BI OR 182177-28-2/BI OR 182177-29-3/BI OR 182177-30-6/BI OR  
182177-31-7/BI OR 182177-32-8/BI OR 182177-33-9/BI OR 182177-34  
-0/BI OR 182177-35-1/BI OR 182177-36-2/BI OR 182177-37-3/BI OR  
182177-38-4/BI OR 182177-39-5/BI OR 182177-40-8/BI OR 182177-41  
-9/BI OR 182177-42-0/BI OR 182177-43-1/BI OR 182177-44-2/BI OR  
182177-45-3/BI OR 182177-46-4/BI OR 182177-47-5/BI OR 182177-48  
-6/BI OR 182177-49-7/BI OR 182177-50-0/BI OR 182177-51-1/BI OR  
182177-52-2/BI OR 182177-53-3/BI OR 182177-54-4/BI OR 182177-55  
-5/BI OR 182177-56-6/BI OR 182177-57-7/BI OR 182177-58-8/BI OR  
182177-59-9/BI OR 182177-60-2/BI OR 182177-61-3/BI OR 182177-62  
-4/BI OR 182177-63-5/BI OR 182177-64-6/BI OR 182177-65-7/BI OR  
182177-66-8/BI OR 182177-67-9/BI OR 182177-68-0/BI OR 182177-69  
-1/BI OR 182177-70-4/BI OR 182177-71-5/BI OR 182177-72-6/BI OR  
182177-73-7/BI OR 182177-74-8/BI OR 182177-75-9/BI OR 182177-76  
-0/BI OR 182177-77-1/BI OR 182177-78-2/BI OR 182177-79-3/BI OR  
182177-80-6/BI OR 182177-81-7/BI OR 182177-82-8/BI OR 182177-83  
-9/BI OR 182177-84-0/BI OR 182177-85-1/BI OR 182177-86-2/BI OR  
182177-87-3/BI OR 182177-88-4/BI OR 182177-89-5/BI OR 182177-90  
-8/BI OR 182177-91-9/BI OR 182177-92-0/BI OR 182177-93-1/BI OR  
182177-94-2/BI OR 182177-95-3/BI OR 182238-37-5/BI OR 182238-38  
-6/BI OR 522671-88-1/BI OR 522671-89-2/BI OR 522672-79-3/BI OR

686373-48-8(BI) AND L24  
D QUE  
SAVE TEMP L86 MOH118SEQ/A

FILE 'STNGUIDE' ENTERED AT 15:55:54 ON 23 FEB 2006

FILE 'CAPLUS' ENTERED AT 15:57:47 ON 23 FEB 2006

D QUE L14  
L87 0 SEA ABB=ON L14 NOT (L83 OR L85)

FILE 'EMBASE' ENTERED AT 15:57:48 ON 23 FEB 2006

D QUE L55  
D QUE L59  
L88 4 SEA ABB=ON (L55 OR L59) NOT L54

FILE 'DRUGU' ENTERED AT 15:57:50 ON 23 FEB 2006

D QUE L71  
L89 0 SEA ABB=ON L71 NOT L62

FILE 'JICST-EPLUS, PASCAL, WPIX, IPA, BIOSIS, ESBIOBASE, BIOTECHDS,  
LIFESCI, CONFSCI, DISSABS, SCISEARCH' ENTERED AT 15:57:52 ON 23 FEB 2006

D QUE L81  
D QUE L82  
L90 6 SEA ABB=ON (L81 OR L82) NOT L80

FILE 'MEDLINE' ENTERED AT 15:57:59 ON 23 FEB 2006

D QUE L42  
D QUE L44  
L91 1 SEA ABB=ON (L42 OR L44) NOT L37

FILE 'STNGUIDE' ENTERED AT 15:58:07 ON 23 FEB 2006

FILE 'MEDLINE, EMBASE, PASCAL, WPIX, BIOSIS, ESBIOBASE, SCISEARCH'  
ENTERED AT 15:58:31 ON 23 FEB 2006

L92 9 DUP REM L91 L88 L90 (2 DUPLICATES REMOVED)  
ANSWER '1' FROM FILE MEDLINE  
ANSWERS '2-5' FROM FILE EMBASE  
ANSWER '6' FROM FILE PASCAL  
ANSWERS '7-8' FROM FILE WPIX  
ANSWER '9' FROM FILE BIOSIS  
D IALL 1-9

FILE 'HOME' ENTERED AT 15:58:46 ON 23 FEB 2006

D SAVED

=>

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=> fil capl; d que 11; d que 15; s 11 or 15; fil medl; d que 137; fil embase; d que 154; fil drugu; d que 162

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FILE COVERS 1907 - 23 Feb 2006 VOL 144 ISS 9  
FILE LAST UPDATED: 22 Feb 2006 (20060222/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

<http://www.cas.org/infopolicy.html>  
'OBI' IS DEFAULT SEARCH FIELD FOR 'CAPLUS' FILE

L1 1 SEA FILE=CAPLUS ABB=ON US2003-660118/AP

*inventor*  
*search*

L3 2951 SEA FILE=CAPLUS ABB=ON WHITE C?/AU  
L4 4538 SEA FILE=CAPLUS ABB=ON THIOREDOXIN#/OBI  
L5 9 SEA FILE=CAPLUS ABB=ON L3 AND L4

L83 9 L1 OR L5

FILE 'MEDLINE' ENTERED AT 15:51:08 ON 23 FEB 2006

FILE LAST UPDATED: 22 FEB 2006 (20060222/UP). FILE COVERS 1950 TO DATE.

On December 11, 2005, the 2006 MeSH terms were loaded.

The MEDLINE reload for 2006 will soon be available. For details on the 2005 reload, enter HELP RLOAD at an arrow prompt (=>). See also:

<http://www.nlm.nih.gov/mesh/>  
[http://www.nlm.nih.gov/pubs/techbull/nd04/nd04\\_mesh.html](http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html)  
[http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\\_med\\_data\\_changes.html](http://www.nlm.nih.gov/pubs/techbull/nd05/nd05_med_data_changes.html)  
[http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\\_2006\\_MeSH.html](http://www.nlm.nih.gov/pubs/techbull/nd05/nd05_2006_MeSH.html)

OLDMEDLINE is covered back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

This file contains CAS Registry Numbers for easy and accurate

L35 2163 SEA FILE=MEDLINE ABB=ON THIOREDOXIN/CT  
L36 2181 SEA FILE=MEDLINE ABB=ON WHITE C?/AU  
L37 7 SEA FILE=MEDLINE ABB=ON L35 AND L36

FILE 'EMBASE' ENTERED AT 15:51:08 ON 23 FEB 2006  
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FILE COVERS 1974 TO 20 Feb 2006 (20060220/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

This file contains CAS Registry Numbers for easy and accurate  
substance identification.

L45 1505 SEA FILE=EMBASE ABB=ON WHITE C?/AU  
L46 2331 SEA FILE=EMBASE ABB=ON THIOREDOXIN/CT  
L54 8 SEA FILE=EMBASE ABB=ON L45 AND L46

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FILE LAST UPDATED: 23 FEB 2006 <20060223/UP>  
>>> DERWENT DRUG FILE (SUBSCRIBER) <<<

>>> FILE COVERS 1983 TO DATE <<<  
>>> THESAURUS AVAILABLE IN /CT <<<

L60 356 SEA FILE=DRUGU ABB=ON WHITE C?/AU  
L61 72 SEA FILE=DRUGU ABB=ON THIOREDOXIN#/CT  
L62 2 SEA FILE=DRUGU ABB=ON L60 AND L61

=> fil jic pascal wpix ipa biosis esbio biotechds lifesci confsci dissabs  
scisearch; d que 180  
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L72 11489 SEA WHITE C?/AU  
L73 17233 SEA THIOREDOXIN#  
L74 38537 SEA SPUTUM  
L75 47183 SEA MUCUS  
L76 3898 SEA MUCOLY?  
L77 74427 SEA LIQUEF?  
L78 567528 SEA VISCO?  
L79 79051 SEA CYSTIC FIBROSIS  
L80 10 SEA L72 AND L73 AND (L74 OR L75 OR L76 OR L77 OR L78 OR L79)

=> dup rem 137,162,183,154,180 >  
FILE 'MEDLINE' ENTERED AT 15:51:39 ON 23 FEB 2006

FILE 'DRUGU' ENTERED AT 15:51:39 ON 23 FEB 2006  
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PROCESSING COMPLETED FOR L37

PROCESSING COMPLETED FOR L62

PROCESSING COMPLETED FOR L83

PROCESSING COMPLETED FOR L54

PROCESSING COMPLETED FOR L80

L84 20 DUP REM L37 L62 L83 L54 L80 (16 DUPLICATES REMOVED)

ANSWERS '1-7' FROM FILE MEDLINE

ANSWERS '8-9' FROM FILE DRUGU

ANSWERS '10-13' FROM FILE CAPLUS

ANSWERS '14-16' FROM FILE EMBASE

ANSWERS '17-18' FROM FILE PASCAL

ANSWERS '19-20' FROM FILE ESBIOBASE

=> d iall 1-9; d ibib ed abs hitind 10-13; d iall 14-20

L84 ANSWER 1 OF 20 MEDLINE on STN DUPLICATE 2  
ACCESSION NUMBER: 2005537349 MEDLINE  
DOCUMENT NUMBER: PubMed ID: 16214824  
TITLE: Thioredoxin and dihydrolipoic acid inhibit elastase activity in cystic fibrosis sputum.  
AUTHOR: Lee Rees L; Rancourt Raymond C; del Val Greg; Pack Kami; Pardee Churee; Accurso Frank J; White Carl W  
CORPORATE SOURCE: Department of Pediatrics, National Jewish Medical and Research Center, Denver, CO 80206, USA.  
CONTRACT NUMBER: HL-07670 (NHLBI)  
SOURCE: American journal of physiology. Lung cellular and molecular physiology, (2005 Nov) 289 (5) L875-82.  
Journal code: 100901229. ISSN: 1040-0605.  
PUB. COUNTRY: United States  
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
LANGUAGE: English  
FILE SEGMENT: Priority Journals  
ENTRY MONTH: 200511  
ENTRY DATE: Entered STN: 20051012  
Last Updated on STN: 20051215  
Entered Medline: 20051129

ABSTRACT:

Excessive neutrophil elastase activity within airways of cystic fibrosis (CF) patients results in progressive lung damage. Disruption of disulfide bonds on elastase by reducing agents may modify its enzymatic activity. Three naturally occurring dithiol reducing systems were examined for their effects on elastase activity: 1) Escherichia coli thioredoxin (Trx) system, 2) recombinant human thioredoxin (rhTrx) system, and 3) dihydrolipoic acid (DHLA). The Trx systems consisted of Trx, Trx reductase, and NADPH. As shown by spectrophotometric assay of elastase activity, the two Trx systems and DHLA inhibited purified human neutrophil elastase as well as the elastolytic activity present in the soluble phase (sol) of CF sputum. Removal of any of the three Trx system constituents prevented inhibition. Compared with the monothiols N-acetylcysteine and reduced glutathione, the dithiols displayed greater elastase inhibition. To streamline Trx as an investigational tool, a stable reduced form of rhTrx was synthesized and used as a single component. Reduced rhTrx inhibited purified elastase and CF sputum sol elastase without NADPH or

Trx reductase. Because Trx and DHLA have mucolytic effects, we investigated changes in elastase activity after mucolytic treatment. Unprocessed CF sputum was directly treated with reduced rhTrx, the Trx system, DHLA, or DNase. The Trx system and DHLA did not increase elastase activity, whereas reduced rhTrx treatment increased sol elastase activity by 60%. By contrast, the elastase activity after DNase treatment increased by 190%. The ability of Trx and DHLA to limit elastase activity combined with their mucolytic effects makes these compounds potential therapies for CF.

CONTROLLED TERM: Adult  
 Animals  
 Child  
 Comparative Study  
 \*Cystic Fibrosis: DT, drug therapy  
 \*Cystic Fibrosis: EN, enzymology  
 Enzyme Inhibitors: PD, pharmacology  
 Escherichia coli Proteins: PD, pharmacology  
 Humans  
 In Vitro  
 \*Leukocyte Elastase: AI, antagonists & inhibitors  
 Rats  
 Recombinant Proteins: PD, pharmacology  
 Research Support, N.I.H., Extramural  
 Research Support, Non-U.S. Gov't  
 Research Support, U.S. Gov't, Non-P.H.S.  
 Research Support, U.S. Gov't, P.H.S.  
 Sputum: EN, enzymology  
 \*Thioctic Acid: AA, analogs & derivatives  
 Thioctic Acid: PD, pharmacology  
 \*Thioredoxin: PD, pharmacology  
 CAS REGISTRY NO.: 462-20-4 (dihydrolipoic acid); 52500-60-4 (Thioredoxin); 62-46-4 (Thioctic Acid)  
 CHEMICAL NAME: 0 (Enzyme Inhibitors); 0 (Escherichia coli Proteins); 0 (Recombinant Proteins); EC 3.4.21.37 (Leukocyte Elastase)

L84 ANSWER 2 OF 20 MEDLINE on STN DUPLICATE 6  
 ACCESSION NUMBER: 2004168104 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 14695120  
 TITLE: Thioredoxin liquefies and decreases the viscoelasticity of cystic fibrosis sputum.  
 AUTHOR: Rancourt Raymond C; Tai Shusheng; King Malcolm; Heltshe Sonya L; Penvari Churee; Accurso Frank J; White Carl W  
 CORPORATE SOURCE: National Jewish Medical and Research Center, 1400 Jackson St., Denver, CO 80206, USA.  
 CONTRACT NUMBER: HL-07670 (NHLBI)  
 SOURCE: American journal of physiology. Lung cellular and molecular physiology, (2004 May) 286 (5) L931-8. Electronic Publication: 2003-12-24.  
 Journal code: 100901229. ISSN: 1040-0605.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Priority Journals  
 ENTRY MONTH: 200405  
 ENTRY DATE: Entered STN: 20040406  
 Last Updated on STN: 20040518  
 Entered Medline: 20040517  
 ABSTRACT:  
 The persistent and viscous nature of airway secretions in cystic fibrosis (CF) disease leads to airway obstruction, opportunistic infection, and deterioration

of lung function. Thioredoxin (Trx) is a protein disulfide reductase that catalyzes numerous thiol-dependent cellular reductive processes. To determine whether Trx can alter the rheological properties of mucus, sputum obtained from CF patients was treated with TRX and its reducing system (0.1 microM thioredoxin reductase + 2 mM NADPH), and liquid phase-gel phase ratio (percent liquid phase) was assessed by compaction assay. Exposure to low Trx concentrations (1 microM) caused significant increases in the percentage of liquid phase of sputum. Maximal increases in percent liquid phase occurred with 30 microM Trx. Additional measurements revealed that sputum liquefaction by the Trx reducing system is dependent on NADPH concentration. The relative potency of the Trx reducing system also was compared with other disulfide-reducing agents. In contrast with Trx, glutathione and N-acetylcysteine were ineffective in liquefying sputum when used at concentrations <1 mM. Sputum viscoelasticity, measured by magnetic microrheometry, also was diminished significantly following 20-min treatment with 3, 10, or 30 microM Trx. Similarly, this reduction in viscoelasticity also was dependent on NADPH concentration. Further investigation has indicated that Trx treatment increases the solubility of high-molecular-weight glycoproteins and causes redistribution of extracellular DNA into the liquid phase of sputum. Recognizing that mucins are the major gel-forming glycoproteins in mucus, we suggest that Trx alters sputum rheology by enzymatic reduction of glycoprotein polymers present in sputum.

CONTROLLED TERM: Check Tags: Female; In Vitro; Male  
 Adolescent  
 Adult  
 Cloning, Molecular  
 \*Cystic Fibrosis: PP, physiopathology  
 Elasticity  
 Escherichia coli  
 Humans  
 Recombinant Proteins: PD, pharmacology  
 Research Support, Non-U.S. Gov't  
 Research Support, U.S. Gov't, P.H.S.  
 Rheology  
 Sputum: DE, drug effects  
 \*Sputum: PH, physiology  
 \*Thioredoxin: PD, pharmacology  
 Viscosity  
 CAS REGISTRY NO.: 52500-60-4 (Thioredoxin)  
 CHEMICAL NAME: O (Recombinant Proteins)

L84 ANSWER 3 OF 20 MEDLINE on STN DUPLICATE 8  
 ACCESSION NUMBER: 1999170595 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 10070119  
 TITLE: Induction of thioredoxin and thioredoxin reductase gene expression in lungs of newborn primates by oxygen.  
 AUTHOR: Das K C; Guo X L; White C W  
 CORPORATE SOURCE: Department of Pediatrics, National Jewish Medical and Research Center, Denver 80206; and University of Colorado Health Sciences Center, Denver, Colorado 80262, USA..  
 kumuda@uthct.edu  
 CONTRACT NUMBER: HL-52732 (NHLBI)  
 HL-53636 (NHLBI)  
 HL-56263 (NHLBI)  
 +  
 SOURCE: American journal of physiology, (1999 Mar) 276 (3 Pt 1)  
 L530-9.  
 Journal code: 0370511. ISSN: 0002-9513.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English  
FILE SEGMENT: Priority Journals  
ENTRY MONTH: 199904  
ENTRY DATE: Entered STN: 19990426  
Last Updated on STN: 19990426  
Entered Medline: 19990415

## ABSTRACT:

Thioredoxin (TRX) is a potent protein disulfide oxidoreductase important in antioxidant defense and regulation of cell growth and signal transduction processes, among them the production of nitric oxide. We report that lung TRX and its reductase, TR, are specifically upregulated at birth by O<sub>2</sub>. Throughout the third trimester, mRNAs for TRX and TR were expressed constitutively at low levels in fetal baboon lungs. However, after premature birth (125 or 140 of 185 days gestation), lung TRX and TR mRNAs increased rapidly with the onset of O<sub>2</sub> or air breathing. Lung TRX mRNA also increased in lungs of term newborns with air breathing. Premature animals (140 days) breathing 100% O<sub>2</sub> develop chronic lung disease within 7-14 days. These animals had greater TRX and TR mRNAs after 1, 6, or 10 days of life than fetal control animals. In 140-day animals given lesser O<sub>2</sub> concentrations (as needed) who do not develop chronic lung disease, lung TRX and TR mRNAs were also increased on days 1 and 6 but not significantly on day 10. In fetal distal lung explant culture, mRNAs for TRX and TR were elevated within 4 h in 95% O<sub>2</sub> relative to 1% O<sub>2</sub>, and the response was similar at various gestations. In contrast, TRX protein did not increase in lung explants from premature animals (125 or 140 days) but did in those from near-term (175-day) fetal baboons after exposure to hyperoxia. However, lung TRX protein and activity, as well as TR activity, eventually did increase in vivo in response to hyperoxia (6 days). Increases in TRX and TR mRNAs in response to 95% O<sub>2</sub> also were observed in adult baboon lung explants. When TRX redox status was determined, increased O<sub>2</sub> tension shifted TRX to its oxidized form. Treatment of lung explants with actinomycin D inhibited TRX and TR mRNA increases in 95% O<sub>2</sub>, indicating transcriptional regulation by O<sub>2</sub>. The acute increase in gene expression for both TRX and TR in response to O<sub>2</sub> suggests an important role for these proteins during the transition from relatively anaerobic fetal life to O<sub>2</sub> breathing at birth.

CONTROLLED TERM: Air  
Animals  
\*Animals, Newborn: PH, physiology  
Culture Techniques  
Delivery, Obstetric  
Fetus: ME, metabolism  
\*Gene Expression Regulation: DE, drug effects  
Gene Expression Regulation: PH, physiology  
Gestational Age  
Humans  
Infant, Newborn  
\*Lung: DE, drug effects  
Lung: EM, embryology  
Lung: ME, metabolism  
Lung: PH, physiology  
\*Oxygen: PD, pharmacology  
Papio  
RNA, Messenger: ME, metabolism  
Research Support, Non-U.S. Gov't  
Research Support, U.S. Gov't, P.H.S.  
Respiration  
Respiratory Distress Syndrome, Newborn: ME, metabolism  
\*Thioredoxin: GE, genetics  
\*Thioredoxin Reductase (NADPH): GE, genetics  
CAS REGISTRY NO.: 52500-60-4 (Thioredoxin); 7782-44-7 (Oxygen)  
CHEMICAL NAME: O (RNA, Messenger); EC 1.6.4.5 (Thioredoxin Reductase)

(NADPH) )

L84 ANSWER 4 OF 20 MEDLINE on STN DUPLICATE 9  
 ACCESSION NUMBER: 1999351908 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 10424622  
 TITLE: Hyperoxia induces thioredoxin and thioredoxin reductase gene expression in lungs of premature baboons with respiratory distress and bronchopulmonary dysplasia.  
 AUTHOR: Das K C; Guo X L; White C W  
 CORPORATE SOURCE: National Jewish Medical and Research Center and University of Colorado Health Sciences Center, Denver 80106, USA.  
 SOURCE: Chest, (1999 Jul) 116 (1 Suppl) 101S.  
 Journal code: 0231335. ISSN: 0012-3692.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals  
 ENTRY MONTH: 199908  
 ENTRY DATE: Entered STN: 19990827  
 Last Updated on STN: 19990827  
 Entered Medline: 19990817  
 CONTROLLED TERM: Animals  
 \*Bronchopulmonary Dysplasia: ME, metabolism  
 Humans  
 Infant, Newborn  
 \*Oxygen: TO, toxicity  
 Papio  
 \*Respiratory Distress Syndrome, Newborn: ME, metabolism  
 \*Thioredoxin: GE, genetics  
 \*Thioredoxin Reductase (NADPH): GE, genetics  
 52500-60-4 (Thioredoxin); 7782-44-7 (Oxygen)  
 EC 1.6.4.5 (Thioredoxin Reductase (NADPH))  
 CAS REGISTRY NO.:  
 CHEMICAL NAME:

L84 ANSWER 5 OF 20 MEDLINE on STN DUPLICATE 10  
 ACCESSION NUMBER: 1998278389 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 9617827  
 TITLE: Detection of thioredoxin in human serum and biological samples using a sensitive sandwich ELISA with digoxigenin-labeled antibody.  
 AUTHOR: Das K C; White C W  
 CORPORATE SOURCE: Department of Pediatrics, National Jewish Medical and Research Center, Denver, CO 80206, USA.  
 CONTRACT NUMBER: HL46481 (NHLBI)  
 HL52732 (NHLBI)  
 HL56263 (NHLBI)  
 SOURCE: Journal of immunological methods, (1998 Feb 1) 211 (1-2) 9-20.  
 Journal code: 1305440. ISSN: 0022-1759.  
 PUB. COUNTRY: Netherlands  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Priority Journals  
 ENTRY MONTH: 199806  
 ENTRY DATE: Entered STN: 19980708  
 Last Updated on STN: 19980708  
 Entered Medline: 19980623

ABSTRACT:  
 Thioredoxin is a low molecular weight, redox active protein important in cellular proliferation, signal transduction and antioxidant function. Thioredoxin is secreted by normal as well as neoplastic cells and is

potentially involved in paracrine cell communication as suggested by its co-cytokine activity. Thus, the thioredoxin level in biological fluids, cells and tissue homogenates could be an important indicator of physiological or pathophysiological conditions. Hence, an accurate and sensitive measurement is of paramount importance in studies involving thioredoxin. We present here an ultrasensitive enzyme linked immuno-absorbent assay (ELISA) for human thioredoxin using digoxigenin-labelled goat polyclonal anti-human thioredoxin. The assay could detect a minimum level of 15 pg/ml thioredoxin in human serum, cell culture media, and in cell and tissue samples. The assay was optimized for concentration of both antibodies, blocking agent, plates, incubation time and reaction volumes. Excellent linearity and reproducibility were obtained. The assay was applied to different baboon tissues and human serum samples. The intrassay coefficient of variation (CV) was between 6.0 to 14 and the interassay CV was from 1.6 to 11.1. Excellent parallelism of standards with serum samples, tissue homogenates or cell lysates was obtained. More than 90% recovery of human thioredoxin was observed in 10% human serum. The assay is easy to use, rapid, reproducible, but above all it is a quantitative, specific and sensitive way to measure thioredoxin in a variety of biological specimens.

CONTROLLED TERM: Animals  
 Antibodies  
 Asthma: BL, blood  
 Buffers  
 Calibration  
 Digoxigenin  
 Dose-Response Relationship, Drug  
 \*Enzyme-Linked Immunosorbent Assay: MT, methods  
 Enzyme-Linked Immunosorbent Assay: ST, standards  
 Goats  
 Horseradish Peroxidase  
 Humans  
 Hydrogen-Ion Concentration  
 Indicators and Reagents  
 Papio: EM, embryology  
 Research Support, Non-U.S. Gov't  
 Research Support, U.S. Gov't, P.H.S.  
 Sensitivity and Specificity  
 \*Thioredoxin: AN, analysis  
 Thioredoxin: BL, blood  
 Time Factors  
 CAS REGISTRY NO.: 1672-46-4 (Digoxigenin); 52500-60-4 (Thioredoxin)  
 CHEMICAL NAME: 0 (Antibodies); 0 (Buffers); 0 (Indicators and Reagents);  
 EC 1.11.1.- (Horseradish Peroxidase)

L84 ANSWER 6 OF 20 MEDLINE on STN DUPLICATE 11  
 ACCESSION NUMBER: 1998072216 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 9409558  
 TITLE: Elevation of manganese superoxide dismutase gene expression by thioredoxin.  
 AUTHOR: Das K C; Lewis-Molock Y; White C W  
 CORPORATE SOURCE: Department of Pediatrics, National Jewish Medical and Research Center, Denver, Colorado 80206, USA.  
 CONTRACT NUMBER: 1R01 HL 52732 (NHLBI)  
 HL46481 (NHLBI)  
 SOURCE: American journal of respiratory cell and molecular biology, (1997 Dec) 17 (6) 713-26.  
 Journal code: 8917225. ISSN: 1044-1549.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Priority Journals

ENTRY MONTH: 199801  
ENTRY DATE: Entered STN: 19980122  
Last Updated on STN: 19980122  
Entered Medline: 19980107

**ABSTRACT:**

Manganese superoxide dismutase (MnSOD) is a mitochondrial enzyme that dismutates potentially toxic superoxide radical into hydrogen peroxide and dioxygen. This enzyme is critical for protection against cellular injury due to elevated partial pressures of oxygen. Thioredoxin (TRX) is a potent protein disulfide reductase found in most organisms that participates in many thiol-dependent cellular reductive processes and plays an important role in antioxidant defense, signal transduction, and regulation of cell growth and proliferation. Here we describe induction of manganese superoxide dismutase by thioredoxin. MnSOD mRNA and activity were increased dramatically by low concentrations of TRX (28 microM). Elevation of MnSOD mRNA by TRX was inhibited by actinomycin D, but not cycloheximide, occurring both in cell lines and primary human lung microvascular endothelial cells. mRNAs for other antioxidant enzymes including copper-zinc superoxide dismutase and catalase were not elevated, demonstrating specificity of induction of MnSOD by TRX. Thiol oxidation by diamide or alkylation by chlorodinitrobenzene inhibited MnSOD induction, further indicating a requirement for reduced TRX. Because both oxidized and reduced thioredoxin (28 microM) induced MnSOD mRNA, the intracellular redox status of externally added Escherichia coli oxidized TRX was determined. About 45% of internalized E. coli TRX was reduced, with 8% in fully reduced form and about 37% in partially reduced form. However, when TRX reductase and nicotinamide adenine dinucleotide (NADPH) were added to the extracellular medium with TRX, more than 80% of E. coli TRX was found to be in a fully reduced state in human adenocarcinoma (A549) cells. Although lower concentrations of oxidized TRX (7 microM) did not induce MnSOD mRNA, this concentration of TRX, when reduced by NADPH and TRX reductase, increased MnSOD mRNA six-fold. In additional studies, MCF-7 cells stably transfected with the human TRX gene had elevated expression of MnSOD mRNA relative to vector-transfected controls. Thus, both endogenously produced and exogenously added TRX elevate MnSOD gene expression. These findings suggest a novel mechanism involving reduced TRX in regulation of MnSOD.

CONTROLLED TERM: Blotting, Western  
Cells, Cultured  
Cycloheximide: PD, pharmacology  
Dactinomycin: PD, pharmacology  
Diamide: PD, pharmacology  
Dinitrochlorobenzene: PD, pharmacology  
Dose-Response Relationship, Drug  
Endothelium, Vascular: DE, drug effects  
Endothelium, Vascular: EN, enzymology  
Enzyme-Linked Immunosorbent Assay  
Escherichia coli: ME, metabolism  
\*Gene Expression Regulation, Enzymologic: DE, drug effects  
Humans  
Kinetics  
Lung: BS, blood supply  
Oxidation-Reduction  
RNA, Messenger: GE, genetics  
Research Support, Non-U.S. Gov't  
Research Support, U.S. Gov't, P.H.S.  
\*Superoxide Dismutase: GE, genetics  
Superoxide Dismutase: ME, metabolism  
\*Thioredoxin: PD, pharmacology  
Tumor Cells, Cultured  
CAS REGISTRY NO.: 10465-78-8 (Diamide); 50-76-0 (Dactinomycin); 52500-60-4 (Thioredoxin); 66-81-9 (Cycloheximide); 97-00-7

CHEMICAL NAME: (Dinitrochlorobenzene)  
 0 (RNA, Messenger); EC 1.15.1.1 (Superoxide Dismutase)

 L84 ANSWER 7 OF 20 MEDLINE on STN  
 ACCESSION NUMBER: 2002389710 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 12122214  
 TITLE: Redox systems of the cell: possible links and implications.  
 COMMENT: Comment on: Proc Natl Acad Sci U S A. 2002 Jul  
 23;99(15):9745-9. PubMed ID: 12119401  
 AUTHOR: Das Kumuda C; White Carl W  
 CORPORATE SOURCE: Department of Molecular Biology, University of Texas at  
 Tyler, 11937 U.S. Highway 271, Tyler, TX 75708, USA.  
 SOURCE: Proceedings of the National Academy of Sciences of the  
 United States of America, (2002 Jul 23) 99 (15) 9617-8.  
 Electronic Publication: 2002-07-16.  
 Journal code: 7505876. ISSN: 0027-8424.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Commentary  
 LANGUAGE: English  
 FILE SEGMENT: Priority Journals  
 ENTRY MONTH: 200209  
 ENTRY DATE: Entered STN: 20020725  
 Last Updated on STN: 20030105  
 Entered Medline: 20020904  
 CONTROLLED TERM: \*Glutathione: ME, metabolism  
 Glyceraldehyde-3-Phosphate Dehydrogenases: ME, metabolism  
 Humans  
 Oxidation-Reduction  
 \*Thioredoxin: ME, metabolism  
 CAS REGISTRY NO.: 52500-60-4 (Thioredoxin); 70-18-8 (Glutathione)  
 CHEMICAL NAME: EC 1.2.1.- (Glyceraldehyde-3-Phosphate Dehydrogenases)

L84 ANSWER 8 OF 20 DRUGU COPYRIGHT 2006 THE THOMSON CORP on STN DUPLICATE 3.  
 ACCESSION NUMBER: 2005-23685 DRUGU B T  
 TITLE: Antioxidant defenses in the preterm lung: role for  
 hypoxia-inducible factors in BPD  
 AUTHOR: Asikainen T M; White C W  
 CORPORATE SOURCE: Nat.Jewish-Med.+Res.Cent.  
 LOCATION: Denver, CO, USA  
 SOURCE: Toxicol.Appl.Pharmacol. (203, No. 2, 177-88, 2005) 3 Fig. 3  
 Tab. 136 Ref.  
 CODEN: TXAPA9 ISSN: 0041-008X  
 AVAIL. OF DOC.: Department of Pediatrics, National Jewish Medical and  
 Research Center, Room D-301, 1400 Jackson Street, Denver, CO  
 80206, U.S.A. (e-mail: asikainent@njc.org).  
 LANGUAGE: English  
 DOCUMENT TYPE: Journal

## ABSTRACT:

Antioxidant defenses in the preterm lung are reviewed. The role for  
 hypoxia-inducible factors in bronchopulmonary dysplasia is discussed.  
 Oxygen-induced lung injury and respiratory distress syndrome and pulmonary  
 antioxidant defenses during development and in hyperoxia are described. The  
 effects of various antioxidants and steroids (classical antioxidant enzymes,  
 extracellular SOD, GSH and thioredoxin peroxidases and their associated  
 reductases GSH and thioredoxin reductases, GSH and thioredoxin, heme  
 oxygenases, and small molecular weight antioxidants (vitamins-C and -E),  
 glucocorticoid, corticosteroids, selenium, inhaled nitric oxide) in preventing

bronchopulmonary dysplasia in preterm neonates are tabulated. This review suggests that single therapeutic factors are insufficient for successful treatment of a preterm baby at risk for developing bronchopulmonary dysplasia.

SECTION HEADING: B Biochemistry  
T Therapeutics

CLASSIF. CODE: 22 Endogenous Compounds  
33 Respiratory  
67 Children and Elderly  
69 Reviews

CONTROLLED TERM:

CASES \*FT; IN-VIVO \*FT; REVIEW \*FT  
[01] BRONCHOPULMONARY \*TR; DYSPNEA \*TR; PNEUMOPATHY \*TR;  
PREMATURE \*FT; INFANT \*FT; MAIN-TOPIC \*FT; ANTIOXIDANT \*FT;  
ANTIOXIDANTS \*FT; PEDIATRICS \*FT; TR \*FT  
[02] ORGOTIEN \*TR; GLUTATHIONE \*TR; THIOREDOXIN \*TR;  
ASCORBATE \*TR; TOCOPHEROL \*TR; SELENIUM-SALT \*TR;  
NITRIC-OXIDE \*TR; TR \*FT

FIELD AVAIL.: AB; LA; CT

FILE SEGMENT: Literature

L84 ANSWER 9 OF 20 DRUGU COPYRIGHT 2006 THE THOMSON CORP on STN DUPLICATE 4  
ACCESSION NUMBER: 2005-41456 DRUGU P B

TITLE: Thioredoxin and dihydrolipoic acid inhibit elastase activity  
in cystic fibrosis sputum.

AUTHOR: Lee R L; Rancourt R C; del Val G; Pack K; Pardee C; Accurso F  
J; White C W

CORPORATE SOURCE: Nat.Jewish-Med.Res.Cent.Denver; Jealot's-Hill.Int.Res.Cent.;  
Univ.Colorado

LOCATION: Denver, CO, USA; Bracknell, U.K.

SOURCE: AJP - Lung Cell.Mol.Physiol. (289, No. 5, L875-L882, 2005) 7  
Fig. 41 Ref. ISSN: 1040-0605

AVAIL. OF DOC.: National Jewish Medical and Research Center, Dept. of  
Pediatrics, Rm. J318, 1400 Jackson St., Denver, CO 80206,  
U.S.A. (C.W.W.). (e-mail: whitec@njc.org).

LANGUAGE: English

DOCUMENT TYPE: Journal

ABSTRACT:

Dihydrolipoic acid (DHLA, dihydrothiostate) and thioredoxin (Trx) are known to decrease the viscoelasticity of cystic fibrosis (CF) mucus. The purpose of this in vitro study was to investigate the effect of DHLA and Trx on elastase activity after mucolytic treatment in CF adult and pediatric patients sputum. Both human (rhTrx) and E. coli recombinant Trx were investigated. Both Trx and DHLA inhibited human neutrophil elastase activity in CF sputum. The level of inhibition was significantly lower for pre-reduced rhTrx compared to rhTrx reduced in situ. A mucolytic effect was shown with pre-reduced rhTrx in whole unprocessed CF sputum but not with DHLA or Trx reduced in situ. The potential therapeutic use of Trx and DHLA, due to their combined elastase and mucolytic effect, in patients with CF is implied.

SECTION HEADING: P Pharmacology  
B Biochemistry

CLASSIF. CODE: 14 Enzyme Inhibitors  
33 Respiratory

## CONTROLLED TERM:

CYSTIC-FIBROSIS \*OC; PNEUMOPATHY \*OC; CONGENITAL-DISEASE \*OC;  
 IN-VITRO \*FT; CASES \*FT; EC-3.4.21.11 \*FT; INHIBITION \*FT;  
 MUCOLYTIC \*FT; SPUTUM \*FT; ELASTASE \*FT

[01] THIOREDOXIN-HUMAN \*PH; THIOREDHU \*RN; RECOMBINANT \*FT; PH \*FT

[02] THIOREDOXIN \*PH; THIOREDOX \*RN; E.COLI \*FT;

RECOMBINANT \*FT; GRAM-NEG. \*FT; BACT. \*FT; PH \*FT

[03] DIHYDROTHIOCTATE \*PH; DIHTHOCT \*RN; ANTIOXIDANTS \*FT; PH \*FT

CAS REGISTRY NO.: 462-20-4

FIELD AVAIL.: AB; LA; CT

FILE SEGMENT: Literature

L84 ANSWER 10 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1

ACCESSION NUMBER: 2005:1103433 CAPLUS

DOCUMENT NUMBER: 143:379832

TITLE: Use of proteins or peptides comprising  
 thioredoxin or lipoic acid as mucolytic and  
 anti-elastase agents for reducing excessively viscous  
 or cohesive mucus or sputum in patients with cystic  
 fibrosis, chronic obstructive pulmonary disease or  
 other disorders

INVENTOR(S): White, Carl W.; Del Val, Greg; Lee, Rees  
 Livingston, II

PATENT ASSIGNEE(S): National Jewish Medical and Research Center, USA;  
 Syngenta Limited; The United States Government

SOURCE: PCT Int. Appl., 123 pp.  
 CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

## PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005094269	A2	20051013	WO 2005-US10061	20050324
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
US 2005260140	A1	20051124	US 2005-90916	20050324
PRIORITY APPLN. INFO.:			US 2004-556516P	P 20040324
			US 2005-650865P	P 20050207
			US 2002-409960P	P 20020910
			US 2003-462082P	P 20030411
			US 2003-660118	A1 20030910

ED Entered STN: 14 Oct 2005

AB The present invention relates to the use of proteins or peptides  
 comprising thioredoxin or lipoic acid as mucolytic and anti-elastase  
 agents for reducing excessively viscous or cohesive mucus or sputum in  
 patients with cystic fibrosis, chronic obstructive pulmonary disease or

other disorders. The compns. contains a compound containing a dithiol active-site in reduced state such as thioredoxin and provides a reducing system for reducing said thioredoxin active site using NADPH and thioredoxin reductase. Respiratory diseases such as CF or COPD are amenable to treatment using compns. described above as well as various gastrointestinal or reproductive disorders.

IC ICM A61K  
 CC 1-9 (Pharmacology)  
 Section cross-reference(s): 63

IT **Thioredoxins**  
 RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (active site, prokaryotic, yeast, plant, mammalian or human; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Drug delivery systems  
 (carriers; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Lung, disease  
 (chronic obstructive pulmonary disease; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Temperature effects, biological  
 (composition administered in absence of elevated; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Peptides, biological studies  
 Proteins  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (containing **thioredoxin** active site; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Thiols, biological studies  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (dithiols, active site, reduced form; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Physiological saline solutions  
 (hypertonic; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Drug delivery systems  
 (inhalants, composition administered via; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Drug delivery systems  
 (intratracheal, composition administered via; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Digestive tract  
 Reproductive system  
 Respiratory system

(mucus in; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Drug delivery systems  
 (nasal, composition administered via; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Drug delivery systems  
 (oral; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Embryophyta  
 Human  
 Mammalia  
 Plant  
 Prokaryota  
 Yeast  
 (**thioredoxin**; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT Cystic fibrosis  
 Mucus  
 Sputum  
 Viscosity  
 (use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 9041-92-3,  $\alpha$ -Antitrypsin 37205-61-1, Proteinase inhibitor  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (composition administered in absence of; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 53-57-6, Nadph 9074-14-0, **Thioredoxin** reductase  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (for reducing **thioredoxin** active site of protein; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 117525-19-6 866665-62-5  
 RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (**thioredoxin** active site sequence; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 866669-22-9 866669-23-0 866669-24-1 866669-25-2 866669-26-3  
 866669-27-4 866669-28-5 866669-29-6 866669-30-9 866669-31-0  
 866669-32-1 866669-33-2  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)  
 (unclaimed protein sequence; use of proteins or peptides comprising **thioredoxin** or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 117525-18-5  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)

(unclaimed sequence; use of proteins or peptides comprising thioredoxin or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 9004-06-2, Elastase  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (use of proteins or peptides comprising thioredoxin or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

IT 70-18-8, Glutathione, biological studies 462-20-4, Dihydrolipoic acid  
 1200-22-2,  $\alpha$ -Lipoic acid 9003-98-9, DNase  
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (use of proteins or peptides comprising thioredoxin or lipoic acid as mucolytic and anti-elastase agents for reducing excessively viscous mucous or sputum in CF or COPD patients)

L84 ANSWER 11 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 5  
 ACCESSION NUMBER: 2004:252597 CAPLUS  
 DOCUMENT NUMBER: 140:281411  
 TITLE: Product and process using a protein or peptide having a thioredoxin active-site in a reduced state for liquefaction of mucus or sputum  
 INVENTOR(S): White, Carl W.  
 PATENT ASSIGNEE(S): National Jewish Medical and Research Center, USA  
 SOURCE: PCT Int. Appl., 69 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 2  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004024868	A2	20040325	WO 2003-US28526	20030910
WO 2004024868	A3	20050519		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2498581	AA	20040325	CA 2003-2498581	20030910
US 2004131606	A1	20040708	US 2003-660118	20030910 <--
EP 1551455	A2	20050713	EP 2003-752262	20030910
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK				
US 2005260140	A1	20051124	US 2005-90916	20050324
PRIORITY APPLN. INFO.:			US 2002-409960P	P 20020910
			US 2003-462082P	P 20030411
			US 2003-660118	A1 20030910
			WO 2003-US28526	W 20030910
			US 2004-556516P	P 20040324
			US 2005-650865P	P 20050207

ED Entered STN: 26 Mar 2004  
 AB The invention discloses compns. and methods for decreasing the viscosity and/or cohesiveness of and/or increasing the liquefaction of excessively

or abnormally viscous or cohesive mucus or sputum. The composition contains a protein or peptide containing a thioredoxin active-site in a reduced state and optionally further contains a reducing system.

IC ICM C12N  
CC 1-12 (Pharmacology)  
Section cross-reference(s): 63  
ST mucus sputum liquefaction protein peptide reduced thioredoxin active site  
IT Drug delivery systems  
(bronchial; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Drug delivery systems  
(direct to lung; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Drug delivery systems  
(inhalants; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Drug delivery systems  
(intratracheal; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Drug delivery systems  
(nasal; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Cystic fibrosis  
Digestive tract  
Drug delivery systems  
Expectorants  
Gastrointestinal agents  
Human  
Lung, disease  
Mucus  
Reproductive system  
Respiratory system  
Sputum  
(protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Thioredoxins  
RL: BSU (Biological study, unclassified); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Proteins  
RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Peptides, biological studies  
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT DNA  
Glycoproteins  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(sputum; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)  
IT Embryophyta  
Escherichia coli  
Mammalia  
Plant

## Prokaryota

## Yeast

(thioredoxin from; protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)

IT 117525-19-6 675625-84-0 675625-85-1  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)

(protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)

IT 70-18-8, Glutathione, biological studies 616-91-1, N-Acetylcysteine 3483-12-3, Dithiothreitol  
 RL: PAC (Pharmacological activity); BIOL (Biological study)  
 (protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)

IT 53-57-6, NADPH 9074-14-0, Thioredoxin reductase  
 RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (protein or peptide with thioredoxin active-site in reduced state for liquefaction of mucus or sputum)

IT 675214-32-1 675214-33-2 675214-34-3 675214-35-4 675214-36-5  
 675214-37-6 675214-38-7 675214-39-8 675214-40-1 675214-41-2  
 675214-42-3 675214-43-4  
 RL: PRP (Properties)  
 (unclaimed protein sequence; product and process using a protein or peptide having a thioredoxin active-site in a reduced state for liquefaction of mucus or sputum)

IT 117525-18-5  
 RL: PRP (Properties)  
 (unclaimed sequence; product and process using a protein or peptide having a thioredoxin active-site in a reduced state for liquefaction of mucus or sputum)

L84 ANSWER 12 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 7

ACCESSION NUMBER: 2002:576282 CAPLUS

DOCUMENT NUMBER: 137:306090

TITLE: Redox systems of the cell: Possible links and implications

AUTHOR(S): Das, Kumuda C.; White, Carl W.

CORPORATE SOURCE: Department of Molecular Biology, University of Texas at Tyler, Tyler, TX, 75708, USA

SOURCE: Proceedings of the National Academy of Sciences of the United States of America (2002), 99(15), 9617-9618

CODEN: PNASA6; ISSN: 0027-8424

PUBLISHER: National Academy of Sciences

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

ED Entered STN: 04 Aug 2002

AB A review discussing a potential link between the two redox systems with glutathione and thioredoxin, and delineating the mechanism by which glutathionylation of thioredoxin can inactivate this multifunctional redox protein. The glutathione and thioredoxin systems are considered parallel redox systems, although their functions are distinct and divergent. Thioredoxin can mediate p53-dependent p21 activation, and thioredoxin translocates from the cytoplasm to the nucleus on stimulation by oxidative stress.

CC 6-0 (General Biochemistry)

ST review redox system glutathione thioredoxin oxidative stress

IT Redox potential

(biol.; role of glutathione and thioredoxin systems in

cellular redox status and oxidative stress)  
 IT Oxidative stress, biological  
     (role of glutathione and thioredoxin systems in cellular  
     redox status and oxidative stress)  
 IT **Thioredoxins**  
   RL: BSU (Biological study, unclassified); BIOL (Biological study)  
     (role of glutathione and thioredoxin systems in cellular  
     redox status and oxidative stress)  
 IT Substitution reaction  
     (thiolation, S-glutathionylation, biol.; role of glutathione and  
     thioredoxin systems in cellular redox status and oxidative  
     stress)  
 IT 70-18-8, Glutathione, biological studies  
   RL: BSU (Biological study, unclassified); BIOL (Biological study)  
     (role of glutathione and thioredoxin systems in cellular  
     redox status and oxidative stress)  
 REFERENCE COUNT: 45 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS  
                   RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 13 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1998:55547 CAPLUS  
 DOCUMENT NUMBER: 128:123821  
 TITLE: Use of thioredoxin-like molecules for  
       induction of manganese-superoxide dismutase (MnSOD) to  
       treat oxidative damage  
 INVENTOR(S): White, Carl W.; Das Kumuda, C.  
 PATENT ASSIGNEE(S): National Jewish Center for Immunology and Respiratory  
                   Medicine, USA  
 SOURCE: PCT Int. Appl., 55 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9800160	A1	19980108	WO 1997-US11167	19970627
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG				
AU 9736434	A1	19980121	AU 1997-36434	19970627
US 5985261	A	19991116	US 1997-883804	19970627
PRIORITY APPLN. INFO.:			US 1996-20740P	P 19960628
			WO 1997-US11167	W 19970627

ED Entered STN: 30 Jan 1998  
 AB A method is provided to increase cellular MnSOD production in an animal to  
   treat oxidative damage; the method involves administering a protein having  
   a thioredoxin active-site in reduced state. A composition and a method to  
   protect an animal from lung disease are provided.  
 IC ICM A61K038-19  
   ICS A61K038-16; A61K038-17  
 CC 1-12 (Pharmacology)  
   Section cross-reference(s): 63  
 ST oxidative damage SOD induction thioredoxin mol; lung disease SOD

induction thioredoxin mol; manganese superoxide dismutase  
induction oxidative damage

IT mRNA  
RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  
(Mn-SOD; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Transcription factors  
RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  
(NF-κB (nuclear factor κB); thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Lung, neoplasm  
(adenocarcinoma; thioredoxin redox status in lung adenocarcinoma cells)

IT Respiratory distress syndrome  
(adult, oxidative damage in; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(bolus; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(capsules; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Kidney  
(cell; thioredoxin effect on Mn-SOD mRNA in different cell types)

IT Surfactants  
(delivery vehicle; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Polymers, biological studies  
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(delivery vehicle; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(diffusion devices; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Lung  
(epithelium, cell; thioredoxin effect on Mn-SOD mRNA in different cell types)

IT Drug delivery systems  
(inhalants; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Reperfusion  
(injury, oxidative damage in; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Lung, disease  
(interstitial, oxidative damage in; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(intratracheal; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(liposomes; thioredoxin-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems

(lipospheres; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(microcapsules; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(microparticles; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Blood vessel  
Blood vessel  
(microvessel, endothelium, cell; **thioredoxin** effect on Mn-SOD mRNA in different cell types)

IT Drug delivery systems  
(nasal; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Respiratory distress syndrome  
(newborn, oxidative damage in; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(oral; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(osmotic pumps; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Asthma  
Atherosclerosis  
Hyperoxia  
Hypoxia, animal  
Inflammation  
Lung, disease  
Neoplasm  
(oxidative damage in; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(parenterals; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Cell  
(recombinant, delivery vehicle; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Drug delivery systems  
(rectal; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Proteins, specific or class  
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(**thioredoxin** active site-containing; **thioredoxin**-like mols. and compns. for induction of manganese-superoxide dismutase to treat oxidative damage)

IT Fibroblast  
(**thioredoxin** effect on Mn-SOD mRNA in different cell types)

IT Redox reaction  
(**thioredoxin** redox status in lung adenocarcinoma cells)

IT Antioxidants  
Drug delivery systems  
Transcription, genetic  
Translation, genetic  
(**thioredoxin**-like mols. and compns. for induction of

manganese-superoxide dismutase to treat oxidative damage)  
 IT Escherichia coli  
 Mammal (Mammalia)  
 Prokaryote  
 Yeast  
 (thioredoxin; thioredoxin-like mols. and compns.  
 for induction of manganese-superoxide dismutase to treat oxidative  
 damage)  
 IT Drug delivery systems  
 (transdermal; thioredoxin-like mols. and compns. for  
 induction of manganese-superoxide dismutase to treat oxidative damage)  
 IT Actins  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 ( $\beta$ -, mRNA; thioredoxin-like mols. and compns. for  
 induction of manganese-superoxide dismutase to treat oxidative damage)  
 IT 9001-05-2, Catalase  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (mRNA; thioredoxin-like mols. and compns. for induction of  
 manganese-superoxide dismutase to treat oxidative damage)  
 IT 9054-89-1, Superoxide dismutase  
 RL: BAC (Biological activity or effector, except adverse); BPR (Biological  
 process); BSU (Biological study, unclassified); THU (Therapeutic use);  
 BIOL (Biological study); PROC (Process); USES (Uses)  
 (manganese-; thioredoxin-like mols. and compns. for induction  
 of manganese-superoxide dismutase to treat oxidative damage)  
 IT 117525-18-5  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL  
 (Biological study)  
 (thioredoxin active site fragment sequence;  
 thioredoxin-like mols. and compns. for induction of  
 manganese-superoxide dismutase to treat oxidative damage)  
 IT 117525-19-6  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL  
 (Biological study)  
 (thioredoxin active site sequence; thioredoxin-like  
 mols. and compns. for induction of manganese-superoxide dismutase to  
 treat oxidative damage)  
 IT 9074-14-0, Thioredoxin reductase  
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological  
 study, unclassified); BIOL (Biological study)  
 (thioredoxin redox status in lung adenocarcinoma cells)  
 REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS  
 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L84 ANSWER 14 OF 20 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights  
 reserved on STM  
 ACCESSION NUMBER: 2004177414 EMBASE  
 TITLE: Thioredoxin liquefies and decreases the viscoelasticity of  
 cystic fibrosis sputum.  
 AUTHOR: Rancourt R.C.; Tai S.; King M.; Heltshe S.L.; Penvari C.;  
 Accurso F.J.; White C.W.  
 CORPORATE SOURCE: C.W. White, Natl. Jewish Med. and Res. Center, 1400 Jackson  
 St., Denver, CO 80206, United States. whitec@njc.org  
 SOURCE: American Journal of Physiology - Lung Cellular and  
 Molecular Physiology, (2004) Vol. 286, No. 5 30-5, pp.  
 L931-L938. .  
 Refs: 35

ISSN: 1040-0605 CODEN: APLPE7  
COUNTRY: United States  
DOCUMENT TYPE: Journal; Article  
FILE SEGMENT: 015 Chest Diseases, Thoracic Surgery and Tuberculosis  
037 Drug Literature Index  
LANGUAGE: English  
SUMMARY LANGUAGE: English  
ENTRY DATE: Entered STN: 20040520  
Last Updated on STN: 20040520

**ABSTRACT:** The persistent and viscous nature of airway secretions in cystic fibrosis (CF) disease leads to airway obstruction, opportunistic infection, and deterioration of lung function. Thioredoxin (Trx) is a protein disulfide reductase that catalyzes numerous thiol-dependent cellular reductive processes. To determine whether Trx can alter the rheological properties of mucus, sputum obtained from CF patients was treated with TRX and its reducing system (0.1  $\mu$ M thioredoxin reductase + 2 mM NADPH), and liquid phase-gel phase ratio (percent liquid phase) was assessed by compaction assay. Exposure to low Trx concentrations (1  $\mu$ M) caused significant increases in the percentage of liquid phase of sputum. Maximal increases in percent liquid phase occurred with 30  $\mu$ M Trx. Additional measurements revealed that sputum liquefaction by the Trx reducing system is dependent on NADPH concentration. The relative potency of the Trx reducing system also was compared with other disulfide-reducing agents. In contrast with Trx, glutathione and N-acetylcysteine were ineffective in liquefying sputum when used at concentrations <1 mM. Sputum viscoelasticity, measured by magnetic microrheometry, also was diminished significantly following 20-min treatment with 3, 10, or 30  $\mu$ M Trx. Similarly, this reduction in viscoelasticity also was dependent on NADPH concentration. Further investigation has indicated that Trx treatment increases the solubility of high-molecular-weight glycoproteins and causes redistribution of extracellular DNA into the liquid phase of sputum. Recognizing that mucins are the major gel-forming glycoproteins in mucus, we suggest that Trx alters sputum rheology by enzymatic reduction of glycoprotein polymers present in sputum.

CONTROLLED TERM: Medical Descriptors:  
\*cystic fibrosis  
\*sputum  
\*liquefaction  
\*viscoelasticity  
airway obstruction: CO, complication  
opportunistic infection: CO, complication  
reduction  
gel  
concentration response  
flow measurement  
mucus  
Western blotting  
DNA content  
solubility  
human  
article  
priority journal  
Drug Descriptors:  
\*thioredoxin  
\*glutathione  
\*acetylcysteine  
\*mucin: EC, endogenous compound  
protein disulfide reductase (glutathione)  
reduced nicotinamide adenine dinucleotide phosphate  
(thioredoxin) 52500-60-4; (glutathione) 70-18-8;

CAS REGISTRY NO.:

(acetylcysteine) 616-91-1; (protein disulfide reductase (glutathione)) 9082-53-5; (reduced nicotinamide adenine dinucleotide phosphate) 53-57-6

COMPANY NAME: American Diagnostica (United States); Sigma (United States); Fisher (United States)

L84 ANSWER 15 OF 20 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights reserved on STN

ACCESSION NUMBER: 2002180321 EMBASE

TITLE: Complete pathway for protein disulfide bond formation encoded by poxviruses.

AUTHOR: Senkevich T.G.; White C.L.; Koonin E.V.; Moss B.

CORPORATE SOURCE: B. Moss, Laboratory of Viral Diseases, Natl. Inst. of Allerg./Infect. Dis., National Institutes of Health, Bethesda, MD 20892, United States. bmoss@nih.gov

SOURCE: Proceedings of the National Academy of Sciences of the United States of America, (14 May 2002) Vol. 99, No. 10, pp. 6667-6672. .

Refs: 27

ISSN: 0027-8424 CODEN: PNASA6

COUNTRY: United States

DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 004 Microbiology

LANGUAGE: English

SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 20020613

Last Updated on STN: 20020613

ABSTRACT: We show that three cytoplasmic thiol oxidoreductases encoded by vaccinia virus comprise a complete pathway for formation of disulfide bonds in intracellular virion membrane proteins. The pathway was defined by analyzing conditional lethal mutants and effects of cysteine to serine substitutions and by trapping disulfide-bonded heterodimer intermediates for each consecutive step. The upstream component, E10R, belongs to the ERV1/ALR family of FAD-containing sulfhydryl oxidases that use oxygen as the electron acceptor. The second component, A2.5L, is a small  $\alpha$ -helical protein with a CxxxxC motif that forms a stable disulfide-linked heterodimer with E10R and a transient disulfide-linked complex with the third component, G4L. The latter is a thioredoxin-like protein that directly oxidizes thiols of L1R, a structural component of the virion membrane with three stable disulfide bonds, and of the related protein F9L. These five proteins are conserved in all poxviruses, suggesting that the pathway is an ancestral mechanism for direct thiol-disulfide interchanges between proteins even in an unfavorable reducing environment.

CONTROLLED TERM: Medical Descriptors:

- \*virus assembly
- protein assembly
- disulfide bond
- Poxvirus
- Vaccinia virus
- virion
- lethal mutant
- amino acid substitution
- electron transport
- alpha helix
- protein structure
- oxidation reduction reaction
- protein expression
- gene overexpression
- covalent bond

nonhuman  
 article  
 priority journal  
 Drug Descriptors:  
 \*thiol derivative  
 \*oxidoreductase  
 \*virus protein  
 \*membrane protein  
 \*flavine adenine nucleotide  
 \*thiol oxidase  
 thioredoxin  
 epitope

CAS REGISTRY NO.: (thiol derivative) 13940-21-1; (oxidoreductase) 9035-73-8,  
 9035-82-9, 9037-80-3, 9055-15-6; (flavine adenine  
 nucleotide) 146-14-5; (thiol oxidase) 9029-39-4;  
 (thioredoxin) 52500-60-4

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 ACCESSION NUMBER: 1999122680 EMBASE  
 TITLE: Induction of thioredoxin and thioredoxin reductase gene  
 expression in lungs of newborn primates by oxygen.  
 AUTHOR: Das K.C.; Guo X.-L.; White C.W.  
 CORPORATE SOURCE: K.C. Das, Dept. of Molecular Biology, Univ. of Texas Health  
 Center, 11937 US Highway 271, Tyler, TX 75708-3154, United  
 States. kumuda@uthct.edu  
 SOURCE: American Journal of Physiology - Lung Cellular and  
 Molecular Physiology, (1999) Vol. 276, No. 3 20-3, pp.  
 L530-L539. .  
 Refs: 50  
 ISSN: 1040-0605 CODEN: APLPE7  
 COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article  
 FILE SEGMENT: 002 Physiology  
 021 Developmental Biology and Teratology  
 029 Clinical Biochemistry  
 LANGUAGE: English  
 SUMMARY LANGUAGE: English  
 ENTRY DATE: Entered STN: 19990429  
 Last Updated on STN: 19990429

ABSTRACT: Thioredoxin (TRX) is a potent protein disulfide oxidoreductase  
 important in antioxidant defense and regulation of cell growth and signal  
 transduction processes, among them the production of nitric oxide. We report  
 that lung TRX and its reductase, TR, are specifically upregulated at birth by  
 O<sub>2</sub>. Throughout the third trimester, mRNAs for TRX and TR were expressed  
 constitutively at low levels in fetal baboon lungs. However, after premature  
 birth (125 or 140 of 185 days gestation), lung TRX and TR mRNAs increased  
 rapidly with the onset of O<sub>2</sub> or air breathing. Lung TRX mRNA also increased in  
 lungs of term newborns with air breathing. Premature animals (140 days)  
 breathing 100% O<sub>2</sub> develop chronic lung disease within 7-14 days. These animals  
 had greater TRX and TR mRNAs after 1, 6, or 10 days of life than fetal control  
 animals. In 140-day animals given lesser O<sub>2</sub> concentrations (as needed) who do  
 not develop chronic lung disease, lung TRX and TR mRNAs were also increased on  
 days 1 and 6 but not significantly on day 10. In fetal distal lung explant  
 culture, mRNAs for TRX and TR were elevated within 4 h in 95% O<sub>2</sub> relative to 1%  
 O<sub>2</sub>, and the response was similar at various gestations. In contrast, TRX  
 protein did not increase in lung explants from premature animals (125 or 140  
 days) but did in those from near-term (175- day) fetal baboons after exposure  
 to hyperoxia. However, lung TRX protein and activity, as well as TR activity,  
 eventually did increase in vivo in response to hyperoxia (6 days). Increases

in TRX and TR mRNAs in response to 95% O<sub>2</sub> also were observed in adult baboon lung explants. When TRX redox status was determined, increased O<sub>2</sub> tension shifted TRX to its oxidized form. Treatment of lung explants with actinomycin D inhibited TRX and TR mRNA increases in 95% O<sub>2</sub>, indicating transcriptional regulation by O<sub>2</sub>. The acute increase in gene expression for both TRX and TR in response to O<sub>2</sub> suggests an important role for these proteins during the transition from relatively anaerobic fetal life to O<sub>2</sub> breathing at birth.

## CONTROLLED TERM: Medical Descriptors:

- \*gene expression
- \*oxygen breathing
- \*fetus lung maturation
- primate
- fetus lung
- protein expression
- antioxidant activity
- cell growth
- signal transduction
- gene expression regulation
- newborn period
- prematurity
- oxygen concentration
- hyperoxia
- lung dysplasia
- respiratory distress
- lung alveolus oxygen tension
- nonhuman
- animal experiment
- controlled study
- animal tissue
- article
- priority journal

## Drug Descriptors:

- \*thioredoxin: EC, endogenous compound
- \*thioredoxin reductase: EC, endogenous compound
- \*oxygen
- nitric oxide: EC, endogenous compound

CAS REGISTRY NO.: (thioredoxin) 52500-60-4; (thioredoxin reductase) 9074-14-0; (oxygen) 7782-44-7; (nitric oxide) 10102-43-9

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ACCESSION NUMBER: 2005-0473450 PASCAL  
COPYRIGHT NOTICE: Copyright .COPYRGT. 2005 INIST-CNRS. All rights reserved.  
TITLE (IN ENGLISH): **Thioredoxin** and dihydrolipoic acid inhibit elastase activity in **cystic fibrosis sputum**  
AUTHOR: LEE Rees L.; RANCOURT Raymond C.; DEL VAL Greg; PACK Kami; PARDEE Churee; ACCURSO Frank J.; **WHITE Carl W.**  
CORPORATE SOURCE: Department of Pediatrics, National Jewish Medical and Research Center, Denver, Colorado, United States; Mike McMorris Cystic Fibrosis Center, University of Colorado Health Sciences Center, Denver, Colorado, United States; Jealott's Hill International Research Center, Bracknell, United Kingdom  
SOURCE: American journal of physiology. Lung cellular and molecular physiology, (2005), 33(5), L875-L882, 41 refs.

DOCUMENT TYPE: ISSN: 1040-0605 CODEN: APLPE7  
BIBLIOGRAPHIC LEVEL: Journal  
COUNTRY: Analytic  
LANGUAGE: United States  
AVAILABILITY: English  
ABSTRACT: INIST-22200, 354000135637220210  
Excessive neutrophil elastase activity within airways of **cystic fibrosis** (CF) patients results in progressive lung damage. Disruption of disulfide bonds on elastase by reducing agents may modify its enzymatic activity. Three naturally occurring dithiol reducing systems were examined for their effects on elastase activity: 1) *Escherichia coli thioredoxin* (Trx) system, 2) recombinant human **thioredoxin** (rhTrx) system, and 3) dihydrolipoic acid (DHLA). The Trx systems consisted of Trx, Trx reductase, and NADPH. As shown by spectrophotometric assay of elastase activity, the two Trx systems and DHLA inhibited purified human neutrophil elastase as well as the elastolytic activity present in the soluble phase (sol) of CF **sputum**. Removal of any of the three Trx system constituents prevented inhibition. Compared with the monothiols N-acetyl-cysteine and reduced glutathione, the dithiols displayed greater elastase inhibition. To streamline Trx as an investigational tool, a stable reduced form of rhTrx was synthesized and used as a single component. Reduced rhTrx inhibited purified elastase and CF **sputum** sol elastase without NADPH or Trx reductase. Because Trx and DHLA have **mucolytic** effects, we investigated changes in elastase activity after **mucolytic** treatment. Unprocessed CF **sputum** was directly treated with reduced rhTrx, the Trx system, DHLA, or DNase. The Trx system and DHLA did not increase elastase activity, whereas reduced rhTrx treatment increased sol elastase activity by 60%. By contrast, the elastase activity after DNase treatment increased by 190%. The ability of Trx and DHLA to limit elastase activity combined with their **mucolytic** effects makes these compounds potential therapies for CF.  
CLASSIFICATION CODE: 002A20; Life sciences; Biological sciences; Vertebrates physiology, Respiratory system  
002B22D05; Life sciences; Medical sciences; Metabolic diseases  
CONTROLLED TERM: **Thioredoxin; Cystic fibrosis; Sputum; Serine endopeptidases; Human; Mammalia; Respiratory system Peptidases; Hydrolases; Enzyme; Vertebrata; Digestive diseases; Respiratory disease; Genetic disease; Metabolic diseases; Pancreatic disease**  
BROADER TERM:  
L84 ANSWER 18 OF 20 PASCAL COPYRIGHT 2006 INIST-CNRS. ALL RIGHTS RESERVED.  
on STN  
ACCESSION NUMBER: 2004-0555882 PASCAL  
COPYRIGHT NOTICE: Copyright .COPYRGT. 2004 INIST-CNRS. All rights reserved.  
TITLE (IN ENGLISH): **Thioredoxin liquefies and decreases the viscoelasticity of**

AUTHOR: **cystic fibrosis sputum**  
RANCOURT Raymond C.; SHUSHENG TAI; KING Malcolm;  
HELTSHY Sonya L.; PENVARI Cheree; ACCURSO Frank J.;  
WHITE Carl W.

CORPORATE SOURCE: Department of Pediatrics, National Jewish Medical and Research Center, Denver 80206, United States; The Mike McMorris Cystic Fibrosis Research and Treatment Center, Department of Pediatrics, University of Colorado School of Medicine, Denver, Colorado 80218, United States; The Children's Hospital, Denver, Colorado 80218, United States; Pulmonary Research Group, University of Alberta, Edmonton, T6G 2S2, Canada

SOURCE: American journal of physiology. Lung cellular and molecular physiology, (2004), 30(5), L931-L938, 35 refs.

ISSN: 1040-0605 CODEN: APLPE7

DOCUMENT TYPE: Journal

BIBLIOGRAPHIC LEVEL: Analytic

COUNTRY: United States

LANGUAGE: English

AVAILABILITY: INIST-22200, 354000111659420050

ABSTRACT: The persistent and **viscous** nature of airway secretions in **cystic fibrosis** (CF) disease leads to airway obstruction, opportunistic infection, and deterioration of lung function. **Thioredoxin** (Trx) is a protein disulfide reductase that catalyzes numerous thiol-dependent cellular reductive processes. To determine whether Trx can alter the rheological properties of **mucus**, **sputum** obtained from CF patients was treated with TRX and its reducing system (0.1  $\mu$ M **thioredoxin** reductase + 2 mM NADPH), and liquid phase-gel phase ratio (percent liquid phase) was assessed by compaction assay. Exposure to low Trx concentrations (1  $\mu$ M) caused significant increases in the percentage of liquid phase of **sputum**. Maximal increases in percent liquid phase occurred with 30  $\mu$ M Trx. Additional measurements revealed that **sputum liquefaction** by the Trx reducing system is dependent on NADPH concentration. The relative potency of the Trx reducing system also was compared with other disulfide-reducing agents. In contrast with Trx, glutathione and N-acetylcysteine were ineffective in liquefying **sputum** when used at concentrations <1 mM. **Sputum viscoelasticity**, measured by magnetic microrheometry, also was diminished significantly following 20-min treatment with 3, 10, or 30  $\mu$ M Trx. Similarly, this reduction in **viscoelasticity** also was dependent on NADPH concentration. Further investigation has indicated that Trx treatment increases the solubility of high-molecular-weight glycoproteins and causes redistribution of extracellular DNA into the liquid phase of **sputum**. Recognizing that mucins are the major gel-forming glycoproteins in **mucus**, we suggest that Trx alters **sputum** rheology by enzymatic reduction of glycoprotein polymers

CLASSIFICATION CODE: present in **sputum**.  
 002A20; Life sciences; Biological sciences;  
 Vertebrates physiology, Respiratory system  
 002B13C03; Life sciences; Medical sciences;  
 Gastroenterology, Digestive system

CONTROLLED TERM: **Thioredoxin**; **Viscoelasticity**;  
**Cystic fibrosis**; **Sputum**;  
**Mucin**; **Mucus**; **Glutathione**; **Acetylcysteine**;  
**Mammalia**; **Respiratory system**

BROADER TERM: **Vertebrata**; **Digestive diseases**; **Respiratory disease**;  
**Genetic disease**; **Metabolic diseases**; **Pancreatic disease**; **Thiol**

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ACCESSION NUMBER: 2005278476 ESBIOBASE

TITLE: **Thioredoxin** and dihydrolipoic acid inhibit elastase activity in **cystic fibrosis** **sputum**  
 Lee R.L.; Rancourt R.C.; Val G.D.; Pack K.; Pardee C.;  
 Accurso F.J.; White C.W.

CORPORATE SOURCE: C.W. White, National Jewish Medical and Research Center, Dept. of Pediatrics, 1400 Jackson St., Denver, CO 80206, United States.  
 E-mail: whitec@njc.org

SOURCE: American Journal of Physiology - Lung Cellular and Molecular Physiology, (2005), 289/5 33-5 (L875-L882), 41 reference(s)  
 CODEN: APLPE7 ISSN: 1040-0605

DOCUMENT TYPE: Journal; Article

COUNTRY: United States

LANGUAGE: English

SUMMARY LANGUAGE: English

ABSTRACT: Excessive neutrophil elastase activity within airways of **cystic fibrosis** (CF) patients results in progressive lung damage. Disruption of disulfide bonds on elastase by reducing agents may modify its enzymatic activity. Three naturally occurring dithiol reducing systems were examined for their effects on elastase activity: 1) *Escherichia coli* **thioredoxin** (Trx) system, 2) recombinant human **thioredoxin** (rhTrx) system, and 3) dihydrolipoic acid (DHLA). The Trx systems consisted of Trx, Trx reductase, and NADPH. As shown by spectrophotometric assay of elastase activity, the two Trx systems and DHLA inhibited purified human neutrophil elastase as well as the elastolytic activity present in the soluble phase (sol) of CF **sputum**. Removal of any of the three Trx system constituents prevented inhibition. Compared with the monothiols N-acetyl-cysteine and reduced glutathione, the dithiols displayed greater elastase inhibition. To streamline Trx as an investigational tool, a stable reduced form of rhTrx was synthesized and used as a single component. Reduced rhTrx inhibited purified elastase and CF **sputum** sol elastase without NADPH or Trx reductase. Because Trx and DHLA have **mucolytic** effects, we investigated changes in elastase activity after **mucolytic** treatment. Unprocessed CF

**sputum** was directly treated with reduced rhTrx, the Trx system, DHLA, or DNase. The Trx system and DHLA did not increase elastase activity, whereas reduced rhTrx treatment increased sol elastase activity by 60%. By contrast, the elastase activity after DNase treatment increased by 190%. The ability of Trx and DHLA to limit elastase activity combined with their **mucolytic** effects makes these compounds potential therapies for CF. Copyright .COPYRGT. 2005 the American Physiological Society.

CLASSIFICATION CODE:

SUPPLEMENTARY TERM:

99 General  
Thioctic acid; Serine protease; Lipoic acid; Human **thioredoxin**; **Mucolytic**

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on STN

ACCESSION NUMBER:

2004108892 ESBIOBASE

TITLE:

**Thioredoxin liquefies and decreases the viscoelasticity of cystic fibrosis sputum**

AUTHOR:

Rancourt R.C.; Tai S.; King M.; Heltshe S.L.; Penvari C.; Accurso F.J.; **White C.W.**

CORPORATE SOURCE:

C.W. White, Natl. Jewish Med. and Res. Center, 1400 Jackson St., Denver, CO 80206, United States.

SOURCE:

American Journal of Physiology - Lung Cellular and Molecular Physiology, (2004), 286/5 30-5 (L931-L938), 35 reference(s)

CODEN: APLPE7 ISSN: 1040-0605

DOCUMENT TYPE:

Journal; Article

COUNTRY:

United States

LANGUAGE:

English

SUMMARY LANGUAGE:

English

ABSTRACT:

The persistent and **viscous** nature of airway secretions in **cystic fibrosis** (CF) disease leads to airway obstruction, opportunistic infection, and deterioration of lung function. **Thioredoxin** (Trx) is a protein disulfide reductase that catalyzes numerous thiol-dependent cellular reductive processes. To determine whether Trx can alter the rheological properties of **mucus**, **sputum** obtained from CF patients was treated with TRX and its reducing system (0.1  $\mu$ M **thioredoxin** reductase + 2 mM NADPH), and liquid phase-gel phase ratio (percent liquid phase) was assessed by compaction assay. Exposure to low Trx concentrations (1  $\mu$ M) caused significant increases in the percentage of liquid phase of **sputum**. Maximal increases in percent liquid phase occurred with 30  $\mu$ M Trx. Additional measurements revealed that **sputum liquefaction** by the Trx reducing system is dependent on NADPH concentration. The relative potency of the Trx reducing system also was compared with other disulfide-reducing agents. In contrast with Trx, glutathione and N-acetylcysteine were ineffective in **liquefying sputum** when used at concentrations <1 mM. **Sputum viscoelasticity**, measured by magnetic microrheometry, also was diminished significantly

following 20-min treatment with 3, 10, or 30  $\mu$ M Trx. Similarly, this reduction in **viscoelasticity** also was dependent on NADPH concentration. Further investigation has indicated that Trx treatment increases the solubility of high-molecular-weight glycoproteins and causes redistribution of extracellular DNA into the liquid phase of **sputum**. Recognizing that mucins are the major gel-forming glycoproteins in **mucus**, we suggest that Trx alters **sputum** rheology by enzymatic reduction of glycoprotein polymers present in **sputum**.

CLASSIFICATION CODE:

SUPPLEMENTARY TERM:

99 General

**Sputum viscoelasticity; Mucin;**  
**Mucus; Glutathione; N-acetylcysteine;**  
**Deoxyribonucleic acid**

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 L18 161373 SEA FILE=CAPLUS ABB=ON VISCO?/OBI  
 L19 1055 SEA FILE=CAPLUS ABB=ON EXPECTORANT#/OBI  
 L24 541003 SEA FILE=REGISTRY ABB=ON .C..C./SQSP  
 L25 271004 SEA FILE=REGISTRY RAN=(,518362-12-4) ABB=ON .C..C./SQSP  
 L26 269999 SEA FILE=REGISTRY ABB=ON L24 NOT L25  
 L27 58456 SEA FILE=CAPLUS ABB=ON L25 OR L26  
 L29 82 SEA FILE=CAPLUS ABB=ON L27 AND (L6 OR L7)  
 L30 8 SEA FILE=CAPLUS ABB=ON L29 AND (L16 OR L17 OR L18 OR L19)

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L85 6 L30 NOT L83 *previously printed w/ inventor search*

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L85 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:402755 CAPLUS

DOCUMENT NUMBER: 140:385998

TITLE: Sputum compaction assay for assessment of respiratory disease therapy

INVENTOR(S): Daugherty, Ann L.; Mrsny, Randy J.; Patapoff, Thomas W.

PATENT ASSIGNEE(S): Genentech, Inc., USA

SOURCE: U.S. Pat. Appl. Publ., 20 pp., Cont. of U.S. Ser. No. 771,078.

CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003165810	A1	20030904	US 2002-162951	20020604
CA 2147469	AA	19940511	CA 1993-2147469	19931102
WO 9410567	A1	19940511	WO 1993-US10519	19931102
W: AU, CA, JP				
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
AU 9455464	A1	19940524	AU 1994-55464	19931102
EP 666985	A1	19950816	EP 1994-900497	19931102
EP 666985	B1	19970716		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE				
JP 08502978	T2	19960402	JP 1993-511391	19931102
AT 155581	E	19970815	AT 1994-900497	19931102
ES 2106493	T3	19971101	ES 1994-900497	19931102
GR 3024987	T3	19980130	GR 1997-402630	19971009

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AU 729265	B2	20010201	AU 1998-60676	19980407
US 2002034727	A1	20020321	US 2001-771078	20010125
US 2005164334	A1	20050728	US 2005-33358	20050110
PRIORITY APPLN. INFO.:				
			US 1992-971019	B1 19921102
			US 1993-132681	B1 19931006
			US 1994-355418	B1 19941213
			US 1995-539468	B1 19951005
			US 1997-840441	B1 19970401
			US 2001-771078	B1 20010125
			AU 1994-55464	A3 19931102
			WO 1993-US10519	W 19931102
			US 2002-162951	A1 20020604

ED Entered STN: 19 May 2004

AB A compaction assay measuring the viscoelasticity of sputum samples of patients subject to respiratory disease is provided. The compaction assay of the present invention is based upon the change in sputum compactability in a centrifugal field following in vitro DNase treatment of sputum, as measured by centrifugal pellet size which is related to the content of large-mol.-weight DNA. This assay is useful in determining the therapeutic efficacy of DNase, antibiotic and other respiratory disease treatments in improving lung function.

IT 686373-48-8

RL: PRP (Properties)  
(unclaimed protein sequence; sputum compaction assay for assessment of respiratory disease therapy)

L85 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:355745 CAPLUS

DOCUMENT NUMBER: 138:364735

TITLE: Characterization, recombinant production and sequence of an acidic mammalian chitinase, and its use in therapy or diagnosis of mucus-associated diseases or infectious diseases

INVENTOR(S): Aerts, Johannes Maria Franciscus Gerardus; Boot, Rolf Gabriel

PATENT ASSIGNEE(S): Neth.

SOURCE: U.S. Pat. Appl. Publ., 27 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003087414	A1	20030508	US 2001-4219	20011102
WO 2003038079	A2	20030508	WO 2002-NL694	20021101
WO 2003038079	A3	20030828		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
EP 1442119	A2	20040804	EP 2002-773037	20021101
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,				

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK				
US 2004253224	A1	20041216	US 2004-787845	20040226
PRIORITY APPLN. INFO.:			US 2001-4219	A 20011102
			WO 2002-NL694	W 20021101

ED Entered STN: 09 May 2003

AB The invention provides a mammalian mucinase capable of hydrolyzing mucin. Cloning, expression, sequences, physicochem. and enzymic properties of human and murine mucinase (acidic mammalian chitinase, AMCase) are described. The mucinase of the invention is among others suitable for counteracting diseases in which mucus is involved. These diseases comprise cystic fibrosis, COPD, asthma, bronchitis, tuberculosis, tumors with altered mucus expression, and mucus-containing pathogens. The invention also provides a pharmaceutical composition comprising an effective amount of

the

mucinase of the invention and a method of therapeutic or prophylactic treatment of an individual against a disease in which mucus is involved. Methods for obtaining the mucinase of the invention are also herewith provided, as well as nucleic acids encoding (part of) the mucinase. In one aspect the invention provides a diagnostic kit comprising a mucinase, a mucinase-specific antibody, a mucinase-derived peptide and/or nucleic acid encoding (part of) said mucinase.

IT 522671-88-1DP, Chitinase (mouse acidic isoenzyme), subfragments are claimed 522671-89-2DP, Chitinase (human acidic isoenzyme), subfragments are claimed

RL: ANT (Analyte); BPN (Biosynthetic preparation); BSU (Biological study, unclassified); COS (Cosmetic use); DGN (Diagnostic use); FFD (Food or feed use); PRP (Properties); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES (Uses)  
(amino acid sequence; characterization, recombinant production and sequence of acidic mammalian chitinase (mucinase), and its use in therapy or diagnosis of mucus-associated diseases or infectious diseases)

IT 522672-79-3

RL: PRP (Properties)  
(unclaimed protein sequence; characterization, recombinant production and sequence of an acidic mammalian chitinase, and its use in therapy or diagnosis of mucus-associated diseases or infectious diseases)

L85 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:122738 CAPLUS

DOCUMENT NUMBER: 136:194272

TITLE: Ribozymes and antisense oligonucleotides for the inhibition of gene expression by calcium-activated chloride channel-1 gene CLCA-1

INVENTOR(S): Thompson, James; McSwiggen, James; McKenzie, Timothy; Ayers, David; Szymkowski, David E.; Grupe, Andrew

PATENT ASSIGNEE(S): Ribozyme Pharmaceuticals, Incorporated, USA; Syntex (U.S.A.) LLC

SOURCE: PCT Int. Appl., 152 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002011674	A2	20020214	WO 2001-US24970	20010809
WO 2002011674	A3	20030925		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,				

GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,  
 LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,  
 RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,  
 UZ, VN, YU, ZA, ZW

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AM, AZ, BY, KG,  
 KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR,  
 IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN,  
 GQ, GW, ML, MR, NE, SN, TD, TG

US 2003064946 A1 20030403 US 2001-927046 20010809

PRIORITY APPLN. INFO.: US 2000-224383P P 20000809

ED Entered STN: 15 Feb 2002

AB Nucleic acid mols., including antisense and enzymic nucleic acid mols., such as hammerhead ribozymes, DNAzymes, and GeneBlocs, which modulate the expression of calcium-activated chloride channels (CLCA1, CLCA2, CLCA3, and CLCA4) are provided. A target discovery target validation approach was used for finding genes that are involved in chronic mucous hypersecretion. The reporter system consists of a plasmid construct, termed pMUC5AC-EGFP, bearing a gene coding for green fluorescent protein (GFP). The promoter region of the GFP gene is replaced by a portion of the mucin 5AC promoter sufficient to direct efficient transcription of the GFP gene; the plasmid also contains the neomycin drug resistance gene. The cell line selected as host for these studies, NCI-H292 (ATCC CRL-1848), is derived from a human lung mucoepidermoid carcinoma. A ribozyme library with two randomized regions comprising six-nucleotide binding "arms" is used to enrich cells for non-responders to mucin induction and a bioinformatics approach used to identify human CLCA1 as a regulator of MUC5AC expression. Antisense, hammerhead, DNAzyme, NCH, amberzyme, zinzyme, and G-Cleaver ribosome binding/cleavage sites in CLCA1 were identified. The nucleic acid mols. are individually analyzed by computer folding to assess whether the sequences fold into the appropriate secondary structure and to anneal to various sites in the RNA target. Those nucleic acid mols. with unfavorable intramol. interactions such as between the binding arms and the catalytic core are eliminated from consideration. Varying binding arm lengths can be chosen to optimize activity.

IT 143831-71-4, Pulmozyme

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (treatment in conjunction with; ribozymes and antisense oligonucleotides for the inhibition of gene expression by calcium-activated chloride channel-1 gene CLCA-1)

L85 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:616381 CAPLUS

DOCUMENT NUMBER: 125:266026

TITLE: Human DNase I variants with low affinity for actin for use in the treatment of respiratory disorders associated with viscous mucus

INVENTOR(S): Lazarus, Robert A.; Shak, Steven; Ulmer, Jana S.

PATENT ASSIGNEE(S): Genentech, Inc., USA

SOURCE: PCT Int. Appl., 148 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9626279	A1	19960829	WO 1996-US2421	19960221
W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE,				

ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK

RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN

WO 9626278 A1 19960829 WO 1995-US2366 19950224

W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MN, MW, MX, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TT

RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG

ES 2188653 T3 20030701 ES 1995-912611 19950224

SK 284191 B6 20041005 SK 1997-1148 19950224

AU 9650263 A1 19960911 AU 1996-50263 19960221

AU 695863 B2 19980827

BR 9607328 A 19971230 BR 1996-7328 19960221

EP 854927 A1 19980729 EP 1996-907094 19960221

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE

PL 184951 B1 20030131 PL 1996-322002 19960221

RU 2215787 C2 20031110 RU 1997-115780 19960221

RO 118886 B1 20031230 RO 1997-1598 19960221

NO 9703877 A 19971024 NO 1997-3877 19970822

PRIORITY APPLN. INFO.: WO 1995-US2366 A 19950224  
US 1995-540527 A 19951010  
EP 1995-912611 A 19950224  
WO 1996-US2421 W 19960221

ED Entered STN: 17 Oct 1996

AB Amino acid substitution variants of human DNase I that have reduced binding affinity for actin are described for use in the treatment of respiratory diseases where problems are associated with high-viscosity mucus, e.g. cystic fibrosis, chronic bronchitis. The variants are obtained by site-directed mutation of the cloned gene and therapeutically effective forms are manufactured by expression of the cloned gene. The invention also relates to pharmaceutical compns. and therapeutic uses of actin-resistant variants of human DNase I.

IT 182177-07-7, Nuclease, deoxyribo-[13-alanine] (human)  
182177-08-8 182177-09-9, Nuclease, deoxyribo-[13-arginine] (human) 182177-10-2 182177-11-3, Nuclease, deoxyribo-[13-tyrosine] (human) 182177-12-4, Nuclease, deoxyribo-[44-alanine] (human) 182177-13-5 182177-14-6, Nuclease, deoxyribo-[44-tyrosine] (human) 182177-15-7 182177-16-8, Nuclease, deoxyribo-[53-alanine] (human) 182177-17-9, Nuclease, deoxyribo-[53-lysine] (human) 182177-18-0, Nuclease, deoxyribo-[53-arginine] (human) 182177-19-1, Nuclease, deoxyribo-[53-tyrosine] (human) 182177-20-4, Nuclease, deoxyribo-[65-alanine] (human) 182177-21-5, Nuclease, deoxyribo-[65-arginine] (human) 182177-22-6 182177-23-7, Nuclease, deoxyribo-[67-alanine] (human) 182177-24-8 182177-25-9, Nuclease, deoxyribo-[67-lysine] (human) 182177-26-0, Nuclease, deoxyribo-[69-lysine] (human) 182177-27-1, Nuclease, deoxyribo-[69-arginine] (human) 182177-28-2 182177-29-3 182177-30-6 182177-31-7 182177-32-8 182177-33-9, Nuclease, deoxyribo-[44-cysteine] (human) 182177-34-0 182177-35-1, Nuclease, deoxyribo-[45-cysteine] (human) 182177-36-2, Nuclease, deoxyribo-[45-lysine] (human) 182177-37-3, Nuclease, deoxyribo-[45-

arginine] (human) 182177-38-4, Nuclease, deoxyribo-[48-cysteine] (human) 182177-39-5, Nuclease, deoxyribo-[48-lysine] (human) 182177-40-8, Nuclease, deoxyribo-[49-cysteine] (human) 182177-41-9 182177-42-0, Nuclease, deoxyribo-[49-arginine] (human) 182177-44-2, Nuclease, deoxyribo-[49-tyrosine] (human) 182177-45-3, Nuclease, deoxyribo-[52-cysteine] (human) 182177-46-4, Nuclease, deoxyribo-[52-lysine] (human) 182177-47-5 182177-48-6, Nuclease, deoxyribo-[53-cysteine] (human) 182177-49-7, Nuclease, deoxyribo-[53-leucine] (human) 182177-50-0 182177-51-1, Nuclease, deoxyribo-[56-cysteine] (human) 182177-52-2 182177-53-3, Nuclease, deoxyribo-[56-lysine] (human) 182177-54-4, Nuclease, deoxyribo-[56-arginine] (human) 182177-55-5 182177-56-6, Nuclease, deoxyribo-[65-cysteine] (human) 182177-57-7, Nuclease, deoxyribo-[65-lysine] (human) 182177-58-8 182177-59-9, Nuclease, deoxyribo-[65-serine] (human) 182177-60-2, Nuclease, deoxyribo-[67-cysteine] (human) 182177-61-3 182177-62-4 182177-63-5 182177-64-6, Nuclease, deoxyribo-[67-proline] (human) 182177-65-7, Nuclease, deoxyribo-[67-arginine] (human) 182177-66-8, Nuclease, deoxyribo-[67-serine] (human) 182177-67-9, Nuclease, deoxyribo-[68-lysine] (human) 182177-68-0 182177-69-1, Nuclease, deoxyribo-[68-arginine] (human) 182177-70-4, Nuclease, deoxyribo-[69-alanine] (human) 182177-71-5, Nuclease, deoxyribo-[69-cysteine] (human) 182177-72-6 182177-73-7 182177-74-8 182177-75-9, Nuclease, deoxyribo-[114-glycine] (human) 182177-76-0 182177-77-1, Nuclease, deoxyribo-[114-lysine] (human) 182177-78-2, Nuclease, deoxyribo-[114-leucine] (human) 182177-79-3 182177-80-6 182177-81-7 182177-82-8 182177-83-9 182177-84-0 182177-85-1 182177-86-2 182177-87-3 182177-88-4 182177-89-5 182177-90-8 182177-91-9 182177-92-0 182177-93-1 182177-94-2, Nuclease, deoxyribo-[48-arginine] (human) 182177-95-3 182238-37-5 182238-38-6, Nuclease, deoxyribo-[65-proline] (human)  
 RL: CAT (Catalyst use); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (amino acid sequence; human DNase I variants with low affinity for actin for use in treatment of respiratory disorders associated with **viscous mucus**)

IT 132053-08-8DP, amino acid-substituted analogs  
 RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)  
 (human DNase I variants with low affinity for actin for use in treatment of respiratory disorders associated with **viscous mucus**)

L85 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN  
 ACCESSION NUMBER: 1996:616378 CAPLUS  
 DOCUMENT NUMBER: 125:257176  
 TITLE: Human DNase I gene was mutated and enzyme was engineered for actin resistance and pharmaceutical use in reducing sputum **viscoelasticity** in lung disease treatment  
 INVENTOR(S): Lazarus, Robert A.; Shak, Steven; Ulmer, Jana S.  
 PATENT ASSIGNEE(S): Genentech, Inc., USA  
 SOURCE: PCT Int. Appl., 38 pp.

CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 2  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9626278	A1	19960829	WO 1995-US2366	19950224
W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MN, MW, MX, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TT				
RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG.				
CA 2210871	AA	19960829	CA 1995-2210871	19950224
AU 9519703	A1	19960911	AU 1995-19703	19950224
AU 720635	B2	20000608		
BR 9510323	A	19971111	BR 1995-10323	19950224
EP 811068	A1	19971210	EP 1995-912611	19950224
EP 811068	B1	20021218		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE				
HU 77422	A2	19980428	HU 1998-180	19950224
JP 11505408	T2	19990521	JP 1995-525640	19950224
PL 180773	B1	20010430	PL 1995-321890	19950224
CZ 288633	B6	20010815	CZ 1997-2677	19950224
AT 230027	E	20030115	AT 1995-912611	19950224
PT 811068	T	20030430	PT 1995-912611	19950224
ES 2188653	T3	20030701	ES 1995-912611	19950224
SK 283850	B6	20040302	SK 1997-1147	19950224
SK 284191	B6	20041005	SK 1997-1148	19950224
CA 2211413	AA	19960829	CA 1996-2211413	19960221
WO 9626279	A1	19960829	WO 1996-US2421	19960221
W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MK, MN, MW, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK				
RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN				
AU 9650263	A1	19960911	AU 1996-50263	19960221
AU 695863	B2	19980827		
BR 9607328	A	19971230	BR 1996-7328	19960221
PL 184951	B1	20030131	PL 1996-322002	19960221
RU 2215787	C2	20031110	RU 1997-115780	19960221
RO 118886	B1	20031230	RO 1997-1598	19960221
IL 117218	A1	20040601	IL 1996-117218	19960221
ZA 9601419	A	19970822	ZA 1996-1419	19960222
BG 64061	B1	20031128	BG 1997-101846	19970821
BG 64423	B1	20050131	BG 1997-101847	19970821
NO 9703876	A	19971024	NO 1997-3876	19970822
NO 9703877	A	19971024	NO 1997-3877	19970822
NZ 334762	A	20001027	NZ 1999-334762	19990322
NZ 505985	A	20020201	NZ 2000-505985	20000726
US 2001041360	A1	20011115	US 2001-796774	20010228
US 6348343	B2	20020219		
PRIORITY APPLN. INFO.:			EP 1995-912611	A 19950224
			WO 1995-US2366	19950224
			US 1995-403873	B2 19950324

US 1995-540527	A 19951010
WO 1996-US2421	W 19960221
US 1997-929995	B1 19970915
NZ 1999-303837	A1 19990322
NZ 2000-282552	A1 20000726

ED Entered STN: 17 Oct 1996

AB The present invention relates to amino acid sequence variants of human DNase I that have reduced binding affinity for actin. The invention provides nucleic acid sequences encoding such actin-resistant variants, thereby enabling the production of these variants in quantities sufficient for clin. use. The invention also relates to pharmaceutical compns. and therapeutic uses of actin-resistant variants of human DNase I. DNase I variants are useful for reducing viscoelasticity of sputum in patients with cystic fibrosis or other pulmonary diseases or disorders.

IT 143831-71-4DP, Nuclease, deoxyribo-(human clone 18-1 protein moiety), mutant derivs.

RL: BPN (Biosynthetic preparation); BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses) (human DNase I gene was mutated and enzyme was engineered for actin resistance and pharmaceutical use in reducing sputum viscoelasticity in lung disease treatment)

L85 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:76410 CAPLUS

DOCUMENT NUMBER: 114:76410

TITLE: Cloning and expression of cDNA for human pancreatic deoxyribonuclease I

INVENTOR(S): Shak, Steven

PATENT ASSIGNEE(S): Genentech, Inc., USA

SOURCE: PCT Int. Appl., 50 pp.  
CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9007572	A1	19900712	WO 1989-US5744	19891220
W: AU, JP RW: AT, BE, CH, DE, ES, FR, GB, IT, LU, NL, SE				
AU 9048265	A1	19900801	AU 1990-48265	19891220
AU 630658	B2	19921105		
EP 449968	A1	19911009	EP 1990-901443	19891220
EP 449968	B1	19990224		
R: AT, BE, CH, DE, ES, FR, GB, IT, LI, LU, NL, SE				
JP 04502406	T2	19920507	JP 1990-501900	19891220
JP 3162372	B2	20010425		
EP 853121	A2	19980715	EP 1998-105190	19891220
EP 853121	A3	19980805		
R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE				
AT 176924	E	19990315	AT 1990-901443	19891220
ES 2130120	T3	19990701	ES 1990-901443	19891220
JP 2001157580	A2	20010612	JP 2000-310722	19891220
CA 2006473	AA	19900623	CA 1989-2006473	19891221
CA 2006473	C	20020205		
US 2003044403	A1	20030306	US 2001-5675	20011107
US 2005009056	A1	20050113	US 2004-839046	20040504
PRIORITY APPLN. INFO.:			US 1988-289958	A 19881223

US 1989-448038	A 19891208
EP 1990-901443	A3 19891220
JP 1990-501900	A3 19891220
WO 1989-US5744	A 19891220
US 1992-914226	B3 19920713
US 1993-117584	B1 19930903
US 1995-528876	B1 19950915
US 1996-761578	B1 19961209
US 2000-669306	B1 20000925
US 2001-5675	B1 20011107

ED Entered STN: 09 Mar 1991

AB A cDNA encoding a human DNase I (DNase I) is cloned and expressed in Escherichia coli and mammalian cell culture. The enzyme is therapeutically useful for lowering the viscosity of sputum, for example in the treatment of cystic fibrosis without causing an immune response to the enzyme. The cDNA was cloned from a pancreatic cDNA library using oligonucleotide probes derived from the amino acid sequence of the bovine enzyme. Expression vectors for E. coli, HEK-293, and CHO cells were constructed using appropriate promoters. Transformants of E. coli with the plasmid pDNA11 (an expression-secretion vector) yielded up to 500 mg DNase I/L. Stable expression of the gene in CHO cells resulted in the manufacture of the enzyme at .apprx.0.05 pg/cell/day. The recombinant enzyme was shown to be capable of lowering the viscosity of sputum from cystic fibrosis patients (qual. determination).

IT 132053-07-7 132053-08-8

RL: PRP (Properties)

(amino acid sequence of and expression in Escherichia coli and animal cell culture of gene for)

=> fil capl; d que 114  
FILE 'CAPLUS' ENTERED AT 15:57:47 ON 23 FEB 2006  
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L7	4370 SEA FILE=CAPLUS ABB=ON	MUCUS/CT
L11	3325 SEA FILE=CAPLUS ABB=ON	THIOREDOXINS/CT
L12	7 SEA FILE=CAPLUS ABB=ON	L11 AND (L6 OR L7)
L13	728128 SEA FILE=CAPLUS ABB=ON	9/SC, SX - <i>Search code - P. B. O'Bryen info</i>
L14	4 SEA FILE=CAPLUS ABB=ON	L12 NOT L13

=> s 114 not (l83 or l85) *new. D.W.W. 4/10/06*

L87 0 L14 NOT (L83 OR L85)

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L48	3562 SEA FILE=EMBASE ABB=ON	SPUTUM/CT
L49	100 SEA FILE=EMBASE ABB=ON	SPUTUM VISCOSITY/CT
L50	3436 SEA FILE=EMBASE ABB=ON	VISCOELASTICITY/CT
L51	9749 SEA FILE=EMBASE ABB=ON	VISCOSITY/CT
L52	267 SEA FILE=EMBASE ABB=ON	MUCOLYSIS/CT
L53	183 SEA FILE=EMBASE ABB=ON	LIQUEFACTION/CT

L55 5 SEA FILE=EMBASE ABB=ON L46 AND (L47 OR L48 OR L49 OR L50 OR  
< L51 OR L52 OR L53)

L46 2331 SEA FILE=EMBASE ABB=ON THIOREDOXIN/CT  
L56 20389 SEA FILE=EMBASE ABB=ON CYSTIC FIBROSIS/CT  
L59 1 SEA FILE=EMBASE ABB=ON L46 (L) DT/CT AND L56

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↑  
subheading DT = drug therapy

L88 4 (L55 OR L59) NOT L54 previously  
=> fil drugu; d que 171 printed

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L63 405 SEA FILE=DRUGU ABB=ON CYSTIC-FIBROSIS/CT  
L64 1059 SEA FILE=DRUGU ABB=ON SPUTUM/CT  
L65 4493 SEA FILE=DRUGU ABB=ON MUCOLYTIC#/CT  
L66 972 SEA FILE=DRUGU ABB=ON MUCUS/CT  
L67 2111 SEA FILE=DRUGU ABB=ON VISCOSITY/CT  
L68 14 SEA FILE=DRUGU ABB=ON LIQUEFACTION/CT OR LIQUEFYING/CT  
L70 45091 SEA FILE=DRUGU ABB=ON RESPIRATORY/CC  
<L71 1 SEA FILE=DRUGU ABB=ON L61 AND (L63 OR L64 OR L65 OR L66 OR  
L67 OR L68) AND L70

=> s 171 not 162

L89 0 L71 NOT L62 previously  
=> fil jic pascal wpix ipa biosis esbio biotechds lifesci confsci dissabs  
scisearch; d que 181; d que 182 printed

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L73 17233 SEA THIOREDOXIN#  
L76 3898 SEA MUCOLY?  
L81 8 SEA L73 AND L76

L73 17233 SEA THIOREDOXIN#  
L74 38537 SEA SPUTUM  
L75 47183 SEA MUCUS  
L77 74427 SEA LIQUEF?  
L78 567528 SEA VISCO?  
L79 79051 SEA CYSTIC FIBROSIS  
L82 14 SEA L73 AND (L74 OR L75) AND (L77 OR L78 OR L79)

=> s 181-182 not 180

L90 6 (L81 OR L82) NOT L80

=> fil medl; d que 142; d que 144

FILE 'MEDLINE' ENTERED AT 15:57:59 ON 23 FEB 2006

FILE LAST UPDATED: 22 FEB 2006 (20060222/UP). FILE COVERS 1950 TO DATE.

On December 11, 2005, the 2006 MeSH terms were loaded.

The MEDLINE reload for 2006 will soon be available. For details on the 2005 reload, enter HELP RLOAD at an arrow prompt (=>).  
See also:

<http://www.nlm.nih.gov/mesh/>  
[http://www.nlm.nih.gov/pubs/techbull/nd04/nd04\\_mesh.html](http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html)  
[http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\\_med\\_data\\_changes.html](http://www.nlm.nih.gov/pubs/techbull/nd05/nd05_med_data_changes.html)  
[http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\\_2006\\_MeSH.html](http://www.nlm.nih.gov/pubs/techbull/nd05/nd05_2006_MeSH.html)

OLDMEDLINE is covered back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

This file contains CAS Registry Numbers for easy and accurate

L35	2163	SEA FILE=MEDLINE ABB=ON	THIOREDOXIN/CT
L38	11646	SEA FILE=MEDLINE ABB=ON	SPUTUM/CT
L39	13832	SEA FILE=MEDLINE ABB=ON	VISCOSITY/CT
L40	8697	SEA FILE=MEDLINE ABB=ON	MUCUS+NT/CT
L42	3	SEA FILE=MEDLINE ABB=ON	L35 AND (L38 OR L39 OR L40)

L35	2163	SEA FILE=MEDLINE ABB=ON	THIOREDOXIN/CT
L43	19620	SEA FILE=MEDLINE ABB=ON	CYSTIC FIBROSIS/CT
L44	2	SEA FILE=MEDLINE ABB=ON	L43 AND L35

=> s (l42 or l44) not l37

L91 1 (L42 OR L44) NOT L37 *previously printed*

=> dup rem 191,188,190  
FILE 'MEDLINE' ENTERED AT 15:58:31 ON 23 FEB 2006

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PROCESSING COMPLETED FOR L91

PROCESSING COMPLETED FOR L88

PROCESSING COMPLETED FOR L90

L92 9 DUP REM L91 L88 L90 (2 DUPLICATES REMOVED)  
ANSWER '1' FROM FILE MEDLINE  
ANSWERS '2-5' FROM FILE EMBASE  
ANSWER '6' FROM FILE PASCAL  
ANSWERS '7-8' FROM FILE WPIX  
ANSWER '9' FROM FILE BIOSIS

=> d iall 1-9; fil hom

L92 ANSWER 1 OF 9 MEDLINE on STN  
 ACCESSION NUMBER: 2002632750 MEDLINE  
 DOCUMENT NUMBER: PubMed ID: 12391249  
 TITLE: A small molecule inhibitor of redox-regulated NF-kappa B and activator protein-1 transcription blocks allergic airway inflammation in a mouse asthma model.  
 AUTHOR: Henderson William R Jr; Chi Emil Y; Teo Jia-Ling; Nguyen Cu; Kahn Michael  
 CORPORATE SOURCE: Department of Medicine, University Washington, Seattle 98195, USA.. joangb@u.washington.edu  
 CONTRACT NUMBER: AI42989 (NIAID)  
 SOURCE: Journal of immunology (Baltimore, Md. : 1950), (2002 Nov 1) 169 (9) 5294-9.  
 PUB. COUNTRY: Journal code: 2985117R. ISSN: 0022-1767.  
 DOCUMENT TYPE: United States  
 LANGUAGE: Journal; Article; (JOURNAL ARTICLE)  
 FILE SEGMENT: English  
 ENTRY MONTH: Abridged Index Medicus Journals; Priority Journals  
 ENTRY DATE: 200212  
 Entered STN: 20021023  
 Last Updated on STN: 20021217  
 Entered Medline: 20021210

**ABSTRACT:**  
 An oxidant/antioxidant imbalance is seen in the lungs of patients with asthma. This oxidative stress in asthmatic airways may lead to activation of redox-sensitive transcription factors, NF-kappaB and AP-1. We examined the effect of the small molecule inhibitor of redox-regulated NF-kappaB and AP-1 transcription, MOL 294 on airway inflammation and airway hyperreactivity (AHR) in a mouse model of asthma. MOL 294 is a potent nonpeptide inhibitor of NF-kappaB and AP-1 based upon a beta-strand template that binds to and inhibits the cellular redox protein thioredoxin. BALB/c mice after i.p. OVA sensitization (day 0) were challenged with intranasal OVA on days 14, 25, 26, and 27. MOL 294, administered intranasal on days 25-27, blocked the airway inflammatory response to OVA assessed 24 h after the last OVA challenge on day 28. MOL 294 reduced eosinophil, IL-13, and eotaxin levels in bronchoalveolar lavage fluid and airway tissue eosinophilia and mucus hypersecretion. MOL 294 also decreased AHR in vivo to methacholine. These results support redox-regulated transcription as a therapeutic target in asthma and demonstrate that selective inhibitors can reduce allergic airway inflammation and AHR.

CONTROLLED TERM: Check Tags: Female  
 Administration, Intranasal  
 \*Allergens: AD, administration & dosage  
 Animals  
 \*Asthma: ME, metabolism  
 Asthma: PA, pathology  
 \*Asthma: PC, prevention & control  
 Bronchial Hyperreactivity: PC, prevention & control  
 Bronchoalveolar Lavage Fluid: CY, cytology  
 Bronchoalveolar Lavage Fluid: IM, immunology  
 Cell Movement: DE, drug effects  
 Cell Movement: IM, immunology  
 Chemokines, CC: BI, biosynthesis  
 Disease Models, Animal  
 Eosinophils: DE, drug effects  
 Eosinophils: PA, pathology  
 Humans  
 Inflammation: ME, metabolism  
 Inflammation: PC, prevention & control  
 Interleukin-13: BI, biosynthesis  
 Lung: DE, drug effects

Lung: IM, immunology  
 \*Lung: PA, pathology  
 Mice  
 Mice, Inbred BALB C  
     Mucus: DE, drug effects  
     Mucus: IM, immunology  
     Mucus: SE, secretion  
 \*NF-kappa B: AI, antagonists & inhibitors  
 NF-kappa B: ME, metabolism  
 Ovalbumin: AD, administration & dosage  
 Ovalbumin: IM, immunology  
 Oxidation-Reduction: DE, drug effects  
 \*Pyridazines: PD, pharmacology  
 Pyridazines: TU, therapeutic use  
 Research Support, U.S. Gov't, P.H.S.  
     Thioredoxin: AI, antagonists & inhibitors  
 \*Transcription Factor AP-1: AI, antagonists & inhibitors  
     Transcription Factor AP-1: ME, metabolism  
 \*Triazoles: PD, pharmacology  
     Triazoles: TU, therapeutic use  
 Tumor Cells, Cultured  
 CAS REGISTRY NO.: 52500-60-4 (Thioredoxin); 9006-59-1 (Ovalbumin)  
 CHEMICAL NAME: 0 (Allergens); 0 (Chemokines, CC); 0 (Interleukin-13); 0 (MOL 294); 0 (NF-kappa B); 0 (Pyridazines); 0 (Transcription Factor AP-1); 0 (Triazoles); 0 (eotaxin)

L92 ANSWER 2 OF 9 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights reserved on STN  
 ACCESSION NUMBER: 2005547205 EMBASE  
 TITLE: Oxidants and COPD.  
 AUTHOR: MacNee W.  
 CORPORATE SOURCE: W. MacNee, ELEGI, Colt Research Laboratories, Medical School, Teviot Place, Edinburgh EH8 9AG, United Kingdom.  
 w.macnee@ed.ac.uk  
 SOURCE: Current Drug Targets: Inflammation and Allergy, (2005) Vol. 4, No. 6, pp. 627-641. .  
 Refs: 162  
 ISSN: 1568-010X CODEN: CDTICU  
 COUNTRY: Netherlands  
 DOCUMENT TYPE: Journal; General Review  
 FILE SEGMENT:  
     004 Microbiology  
     015 Chest Diseases, Thoracic Surgery and Tuberculosis  
     022 Human Genetics  
     026 Immunology, Serology and Transplantation  
     030 Pharmacology  
     037 Drug Literature Index  
 LANGUAGE: English  
 SUMMARY LANGUAGE: English  
 ENTRY DATE: Entered STN: 20051222  
                   Last Updated on STN: 20051222  
 ABSTRACT: Smoking is the main etiologic factor in chronic obstructive pulmonary disease (COPD). Cigarette smoke produces an enormous oxidant burden on the lungs, which is exacerbated by the release of oxidants from inflammatory cells. There is considerable evidence that an increased oxidative burden occurs in the lungs of patients with COPD, and this may be involved in many of the pathogenic processes, such as direct injury to lung cells, mucus hypersecretion, inactivation of antiproteases, and enhancing lung inflammation through activation of redox-sensitive transcription factors. COPD is also recognized to have multiple systemic consequences, such as weight loss and skeletal muscle dysfunction. Moreover, it is appreciated that oxidative stress

extends beyond the lung and may, through similar oxidative stress mechanisms as those in the lung, contribute to several of the systemic manifestations in COPD such as skeletal muscle dysfunction. Thus, there is a great need for an effective antioxidant therapy to modulate the oxidative stress in COPD, since this may be an important therapeutic target. .COPYRGT. 2005 Bentham Science Publishers Ltd.

CONTROLLED TERM: Medical Descriptors:

- \*chronic obstructive lung disease: DT, drug therapy
- \*chronic obstructive lung disease: ET, etiology
- \*chronic obstructive lung disease: PC, prevention
- risk assessment
- risk factor
- cigarette smoking
- air pollution
- dietary intake
- vitamin supplementation
- disease exacerbation
- inflammatory cell
- pathogenesis
- lung alveolus cell
- cell damage
- oxidation reduction reaction
- weight reduction
- body weight disorder: CO, complication
- myopathy: CO, complication
- oxidative stress
- lung biopsy
- pathophysiology
- lymphocyte function
- defense mechanism
- lung alveolus epithelium
- lung alveolus macrophage
- in vitro study
- signal transduction
- forced expiratory volume
- lung function test
- breath analysis
- lung lavage
- lipid peroxidation
- mucus secretion
- bronchus mucus
- gene expression regulation
- gene silencing
- virus infection
- apoptosis
- exercise
- glutathione metabolism
- muscle metabolism
- muscle atrophy: CO, complication
- protein expression
- protein function
- antioxidant activity
- human
- nonhuman
- review

Drug Descriptors:

- \*oxidizing agent
- proteinase inhibitor: EC, endogenous compound
- transcription factor: EC, endogenous compound

reactive oxygen metabolite: EC, endogenous compound  
 reactive nitrogen species: EC, endogenous compound  
 cigarette smoke  
 nitric oxide: EC, endogenous compound  
 ozone  
 tumor necrosis factor alpha: EC, endogenous compound  
 lipopolysaccharide: EC, endogenous compound  
 xanthine dehydrogenase: EC, endogenous compound  
 superoxide: EC, endogenous compound  
 cytochrome P450: EC, endogenous compound  
 reduced nicotinamide adenine dinucleotide phosphate: EC, endogenous compound  
 nitric oxide synthase: EC, endogenous compound  
 aldehyde oxidase: EC, endogenous compound  
 flavoprotein: EC, endogenous compound  
 tryptophan 2,3 dioxygenase: EC, endogenous compound  
 iron  
 lipid peroxide  
 superoxide dismutase: EC, endogenous compound  
 catalase: EC, endogenous compound  
 glutathione: EC, endogenous compound  
 thioredoxin: EC, endogenous compound  
 ascorbic acid: DT, drug therapy  
 ascorbic acid: PD, pharmacology  
 beta carotene: DT, drug therapy  
 beta carotene: PD, pharmacology  
 flavonoid: DT, drug therapy  
 immunoglobulin enhancer binding protein: EC, endogenous compound  
 protein kinase: EC, endogenous compound  
 unindexed drug  
 (proteinase inhibitor) 37205-61-1; (nitric oxide) 10102-43-9; (ozone) 10028-15-6; (xanthine dehydrogenase) 9054-84-6; (superoxide) 11062-77-4; (cytochrome P450) 9035-51-2; (reduced nicotinamide adenine dinucleotide phosphate) 53-57-6; (nitric oxide synthase) 125978-95-2; (aldehyde oxidase) 9029-07-6; (tryptophan 2,3 dioxygenase) 9014-51-1; (iron) 14093-02-8, 53858-86-9, 7439-89-6; (superoxide dismutase) 37294-21-6, 9016-01-7, 9054-89-1; (catalase) 9001-05-2; (glutathione) 70-18-8; (thioredoxin) 52500-60-4; (ascorbic acid) 134-03-2, 15421-15-5, 50-81-7; (beta carotene) 7235-40-7; (protein kinase) 9026-43-1

CAS REGISTRY NO.:

(proteinase inhibitor) 37205-61-1; (nitric oxide) 10102-43-9; (ozone) 10028-15-6; (xanthine dehydrogenase) 9054-84-6; (superoxide) 11062-77-4; (cytochrome P450) 9035-51-2; (reduced nicotinamide adenine dinucleotide phosphate) 53-57-6; (nitric oxide synthase) 125978-95-2; (aldehyde oxidase) 9029-07-6; (tryptophan 2,3 dioxygenase) 9014-51-1; (iron) 14093-02-8, 53858-86-9, 7439-89-6; (superoxide dismutase) 37294-21-6, 9016-01-7, 9054-89-1; (catalase) 9001-05-2; (glutathione) 70-18-8; (thioredoxin) 52500-60-4; (ascorbic acid) 134-03-2, 15421-15-5, 50-81-7; (beta carotene) 7235-40-7; (protein kinase) 9026-43-1

L92 ANSWER 3 OF 9 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights reserved on STN

ACCESSION NUMBER: 2004119423 EMBASE

TITLE: Reduced Spectral Density Mapping of a Partially Folded Fragment of *E. coli* Thioredoxin.

AUTHOR: Daughdrill G.W.; Vise P.D.; Zhou H.; Yang X.; Yu W.-F.; Tasayco M.L.; Lowry D.F.

CORPORATE SOURCE: G.W. Daughdrill, Department of Microbiology, University of Idaho, P.O. Box 443052, Moscow, ID 83844-3052, United States. gdaugh@uidaho.edu

SOURCE: Journal of Biomolecular Structure and Dynamics, (2004) Vol. 21, No. 5, pp. 663-670. .

Refs: 18

ISSN: 0739-1102 CODEN: JBSDD6

COUNTRY: United States

DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 004 Microbiology

LANGUAGE: English  
 SUMMARY LANGUAGE: English  
 ENTRY DATE: Entered STN: 20040325  
 Last Updated on STN: 20040325

ABSTRACT: The backbone dynamics of a partially folded, N-terminal fragment of *E. coli* thioredoxin were investigated using nuclear magnetic resonance spectroscopy (NMR). Relaxation data were collected at three temperatures and analyzed using reduced spectral density mapping. As temperature was increased, the values for the viscosity normalized  $J(0)$  and for  $J(\omega(H))$  increased, while  $J(\omega(N))$  decreased. The global trend observed for the viscosity normalized  $J(0)$  was consistent with an increase in the hydrodynamic volume of the fragment and suggested the presence of correlated rotational motion in the absence of long range interactions. In addition, the residue specific variation observed for the viscosity normalized  $J(0)$  suggested contributions to  $J(\omega)$  from a range of correlation times that are close to the global correlation time.

CONTROLLED TERM: Medical Descriptors:  
 \*spectrometry  
 \*protein folding  
 \*Escherichia coli  
 molecular dynamics  
 amino terminal sequence  
 nuclear magnetic resonance spectroscopy  
 temperature  
 viscosity  
 hydrodynamics  
 rotation  
 protein interaction  
 correlation function  
 nonhuman  
 article  
 priority journal  
 Drug Descriptors:  
 \*thioredoxin  
 CAS REGISTRY NO.: (thioredoxin) 52500-60-4

L92 ANSWER 4 OF 9 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights reserved on STN  
 ACCESSION NUMBER: 2004512110 EMBASE  
 TITLE: The role of pyocyanin in *Pseudomonas aeruginosa* infection.  
 AUTHOR: Lau G.W.; Hassett D.J.; Ran H.; Kong F.  
 CORPORATE SOURCE: gee.lau@uc.edu  
 SOURCE: Trends in Molecular Medicine, (2004) Vol. 10, No. 12, pp. 599-606.  
 Refs: 60  
 ISSN: 1471-4914 CODEN: TMMRCY  
 PUBLISHER IDENT.: S 1471-4914(04)00260-6  
 COUNTRY: United Kingdom  
 DOCUMENT TYPE: Journal; General Review  
 FILE SEGMENT: 005 General Pathology and Pathological Anatomy  
 017 Public Health, Social Medicine and Epidemiology  
 022 Human Genetics  
 030 Pharmacology  
 037 Drug Literature Index  
 LANGUAGE: English  
 SUMMARY LANGUAGE: English  
 ENTRY DATE: Entered STN: 20041217  
 Last Updated on STN: 20041217  
 ABSTRACT: Pyocyanin (PCN) is a blue redox-active secondary metabolite that is

produced by *Pseudomonas aeruginosa*. PCN is readily recovered in large quantities in sputum from patients with cystic fibrosis who are infected by *P. aeruginosa*. Despite in vitro studies demonstrating that PCN interferes with multiple cellular functions, its importance during clinical infection is uncertain. This is partially caused by the difficulty in defining the contribution of PCN among the numerous virulence factors produced by *P. aeruginosa* during infection. In addition, few cellular pathways that are affected by PCN are known. This review briefly highlights recent advances that might clarify the role of PCN in *P. aeruginosa* pathogenesis.

## CONTROLLED TERM: Medical Descriptors:

- \*bacterial infection: DT, drug therapy
- \*bacterial infection: EP, epidemiology
- \*bacterial infection: ET, etiology
  - \*cystic fibrosis: DT, drug therapy
  - \*cystic fibrosis: EP, epidemiology
  - \*cystic fibrosis: ET, etiology
- \*molecular biology
- \*respiratory tract infection: DT, drug therapy
- \*respiratory tract infection: EP, epidemiology
- \*respiratory tract infection: ET, etiology
- Pseudomonas aeruginosa*
- pathogenesis
- correlation analysis
- biosynthesis
- operon
- bacterial genome
- bioaccumulation
- bacterial virulence
- protein synthesis
- signal transduction
- protein induction
- molecular evolution
- genetic code
- transcription initiation
- enzyme activation
- genetic conservation
- genetic variability
- oxidation reduction reaction
- gene targeting
- Caenorhabditis elegans*
- Saccharomyces cerevisiae*
- enzyme inactivation
- mitochondrial respiration
- gene mutation
- protein localization
- gene expression
- pathophysiology
- protein depletion
- oxidative stress
- human
- nonhuman
- review

Drug Descriptors:

- \*pyocyanine
- protein derivative: EC, endogenous compound
- adenosine triphosphatase: EC, endogenous compound
- antioxidant: DT, drug therapy
  - thioredoxin: DT, drug therapy
- thioredoxin: PD, pharmacology

CAS REGISTRY NO.: (pyocyanine) 85-66-5; (adenosine triphosphatase) 37289-25-1, 9000-83-3; (thioredoxin) 52500-60-4

L92 ANSWER 5 OF 9 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights reserved on STN  
ACCESSION NUMBER: 2004465508 EMBASE  
TITLE: Potential for antioxidant therapy of cystic fibrosis.  
AUTHOR: Cantin A.M.  
CORPORATE SOURCE: A.M. Cantin, Pulmonary Division, Dept. of Med. Faculty of Medicine, University of Sherbrooke, 3001, 12e Avenue Nord, Sherbrooke, Que. J1H 5N4, Canada.  
andre.cantin@usherbrooke.ca  
SOURCE: Current Opinion in Pulmonary Medicine, (2004) Vol. 10, No. 6, pp. 531-536. .  
Refs: 42  
ISSN: 1070-5287 CODEN: COPMFY  
COUNTRY: United States  
DOCUMENT TYPE: Journal; General Review  
FILE SEGMENT: 015 Chest Diseases, Thoracic Surgery and Tuberculosis  
029 Clinical Biochemistry  
030 Pharmacology  
037 Drug Literature Index  
038 Adverse Reactions Titles  
LANGUAGE: English  
SUMMARY LANGUAGE: English  
ENTRY DATE: Entered STN: 20041129  
Last Updated on STN: 20041129  
ABSTRACT: Purpose of review: Changes in redox state clearly play a role in airway inflammation and mucus rheology. Furthermore CFTR (cystic fibrosis transmembrane conductance regulator), the defective protein in cystic fibrosis (CF), not only is regulated by redox state but also directly modulates the epithelial redox environment through transepithelial flux of glutathione. The purpose of this review is to explore the potential therapeutic interest of antioxidant molecules in CF. Recent findings: Several antioxidants have been shown to have mucolytic and anti-inflammatory properties. Some antioxidants such as zinc and vitamin C may also help increase epithelial chloride secretion through CFTR-dependent and independent pathways. Other antioxidants are showing promise in helping CFTR mobilization to plasma membranes. Summary: The many levels of potential application offered by antioxidants make this class of molecules one of the promising areas of therapeutic development for CF. Several redox-modulating agents have a high likelihood of providing useful approaches for the treatment of many aspects of CF airway disease.

CONTROLLED TERM: Medical Descriptors:  
\*cystic fibrosis: DT, drug therapy  
oxidation reduction reaction  
inflammation  
mucus  
lung infection  
diet  
chloride channel  
genetic transcription  
oxidative stress  
gene expression  
respiratory tract disease: SI, side effect  
human  
nonhuman  
review  
Drug Descriptors:  
\*antioxidant

\*zinc: CB, drug combination  
 \*zinc: PD, pharmacology  
 \*ascorbic acid: PD, pharmacology  
 transmembrane conductance regulator: EC, endogenous compound  
 glutathione: DO, drug dose  
 glutathione: DT, drug therapy  
 glutathione: EC, endogenous compound  
 glutathione: PR, pharmaceutics  
 glutathione: IH, inhalational drug administration  
 chloride: EC, endogenous compound  
 reactive oxygen metabolite  
 nacystelyn: AE, adverse drug reaction  
 nacystelyn: DT, drug therapy  
 nacystelyn: PD, pharmacology  
 acetylcysteine: AE, adverse drug reaction  
 acetylcysteine: DT, drug therapy  
 acetylcysteine: PD, pharmacology  
 thioredoxin: EC, endogenous compound  
 reduced nicotinamide adenine dinucleotide phosphate  
 adenosine triphosphate  
 taurine  
 s nitrosoglutathione  
 curcumin: PD, pharmacology  
 selenocystine  
 glutathione derivative  
 alpha tocopherol succinate  
 alpha tocopherylquinone  
 thioctic acid  
 alpha tocopherol  
 immunoglobulin enhancer binding protein: EC, endogenous compound  
 protein kinase C: EC, endogenous compound  
 mucin: EC, endogenous compound  
 glutathione peroxidase: EC, endogenous compound  
 calnexin: EC, endogenous compound  
 calreticulin: EC, endogenous compound  
 epidermal growth factor receptor: EC, endogenous compound  
 toll like receptor 4: EC, endogenous compound  
 unindexed drug  
 unclassified drug

CAS REGISTRY NO.: (zinc) 7440-66-6; (ascorbic acid) 134-03-2, 15421-15-5, 50-81-7; (glutathione) 70-18-8; (chloride) 16887-00-6; (acetylcysteine) 616-91-1; (thioredoxin) 52500-60-4; (reduced nicotinamide adenine dinucleotide phosphate) 53-57-6; (adenosine triphosphate) 15237-44-2, 56-65-5, 987-65-5; (taurine) 107-35-7; (s nitrosoglutathione) 57564-91-7; (curcumin) 458-37-7; (selenocystine) 1464-43-3, 2897-21-4, 29621-88-3; (alpha tocopherol succinate) 17407-37-3, 4345-03-3; (alpha tocopherylquinone) 7559-04-8; (thioctic acid) 1077-29-8, 1200-22-2, 2319-84-8, 62-46-4; (alpha tocopherol) 1406-18-4, 1406-70-8, 52225-20-4, 58-95-7, 59-02-9; (protein kinase C) 141436-78-4; (glutathione peroxidase) 9013-66-5; (calnexin) 139873-08-8; (toll like receptor 4) 203811-83-0

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 ACCESSION NUMBER: 2004-0411858 PASCAL  
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reserved.

TITLE (IN ENGLISH): Possible (enzymatic) routes and biological sites for metabolic reduction of BNP7787, a new protector against cisplatin-induced side-effects

AUTHOR: VERSCHRAAGEN Miranda; BOVEN Epie; TORUN Emine; HAUSHEER Frederick H.; BASF Aalt; VAN DER VIJGH Wim J. F.

CORPORATE SOURCE: Department of Medical Oncology, Vrije Universiteit medical center, De Boelelaan 1117, 1007MB Amsterdam, Netherlands; BioNumerik Pharmaceuticals, Inc., Ste. 400, 8122 Datapoint Drive, San Antonio, TX 78229, United States; Department of Pharmacology and Toxicology, University of Maastricht, P.O. Box 616, 6200MD Maastricht, Netherlands

SOURCE: Biochemical pharmacology, (2004), 68(3), 493-502, 22 refs.

ISSN: 0006-2952 CODEN: BCPCA6

DOCUMENT TYPE: Journal

BIBLIOGRAPHIC LEVEL: Analytic

COUNTRY: United States

LANGUAGE: English

AVAILABILITY: INIST-1418, 354000113766710100

ABSTRACT: Disodium 2,2'-dithio-bis-ethane sulfonate (BNP7787) is under investigation as a potential new chemoprotector against cisplatin-induced nephrotoxicity. The selective protection of BNP7787 appears to arise from the preferential uptake of the drug in the kidneys, where BNP7787 would undergo intracellular conversion into mesna (2-mercapto ethane sulfonate), which in turn can prevent cisplatin induced toxicities. In the present study, we have investigated whether the reduction of BNP7787 into the reactive compound mesna is restricted to the kidney or whether it can also occur in other organs, cells and physiological compartments, including the cytosolic fraction of the renal cortex, plasma, red blood cells (RBCs), liver and small intestine from rats and several tumors (OVCAR-3, MRI-H-207 and WARD). We also determined whether the endogenous thiols glutathione (GSH) and cysteine and the enzyme systems glutaredoxin and thioredoxin, which are all present in the kidney, can be involved in the BNP7787 reduction. UV detection and micro-HPLC with dual electrochemical detection were used to analyze the various incubation mixtures. Our observations are that, in contrast to plasma, a very large reductive conversion of BNP7787 to mesna was measured in RBC lysate. Intact RBCs, however, did not take up BNP7787. Although BNP7787 could be reduced in cytosol of liver and several tumors, this reduction will not be relevant in vivo, since these tissues do not take up large amounts of BNP7787. Kidney cortex cytosol was, similar to the small intestine cytosol, able to substantially reduce BNP7787 to mesna. The ability to reduce BNP7787 in the presence of the endogenous thiols GSH and cysteine, the glutaredoxin system as well as the thioredoxin system, could at least in part explain the high BNP7787 reductive activity of the kidney cortex cytosol. In conclusion, the high reduction of BNP7787 into mesna in the kidney as well

as our earlier observation that the distribution of BNP7787 and mesna was mainly restricted to rat kidney are strong arguments in favor of selective protection of the kidney by BNP7787.

CLASSIFICATION CODE: 002B02; Life sciences; Medical sciences; Pharmacology  
 CONTROLLED TERM: Cisplatin; Toxicity; Mesna; Kidney;  
 Thioredoxin; Pharmacology; Antineoplastic agent; Mucolytic; Uroprotective agent  
 BROADER TERM: Alkylating agent; Urinary system

L92 ANSWER 7 OF 9 WPIX COPYRIGHT 2006 THE THOMSON CORP on STN  
 ACCESSION NUMBER: 2006-079352 [08] WPIX  
 DOC. NO. NON-CPI: N2006-068753  
 DOC. NO. CPI: C2006-028699  
 TITLE: Diagnosing Pseudomonas aeruginosa infection in a subject by detecting in a biological sample from the subject a protein of Pseudomonas aeruginosa, or its modified form, immunogenic fragment or epitope or antibody.  
 DERWENT CLASS: B04 D16 S03  
 INVENTOR(S): PEDERSEN, S K; SLOANE, A J; WEINBERGER, R  
 PATENT ASSIGNEE(S): (PROT-N) PROTEOME SYSTEMS INTELLECTUAL PROPERTY P  
 COUNTRY COUNT: 111  
 PATENT INFORMATION:

PATENT NO	KIND	DATE	WEEK	LA	PG	MAIN	IPC
WO 2006000056	A1	20060105 (200608)*	EN	103	G01N033-50		
RW:	AT BE BG BW CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IS IT KE LS LT LU MC MW MZ NA NL OA PL PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW						
W:	AE AG AL AM AT AU AZ BA BB BG BR BW BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE EG ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KM KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NA NG NI NO NZ OM PG PH PL PT RO RU SC SD SE SG SK SL SM SY TJ TM TN TR TT TZ UA UG US UZ VC VN YU ZA ZM ZW						

## APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 2006000056	A1	WO 2005-AU942	20050628

PRIORITY APPLN. INFO: AU 2004-903521 20040628

## INT. PATENT CLASSIF.:

MAIN: G01N033-50

SECONDARY: A61K039-104; A61K039-40; G01N033-53; G01N033-68

## BASIC ABSTRACT:

WO2006000056 A UPAB: 20060201

NOVELTY - Diagnosing an infection caused by Pseudomonas aeruginosa in a subject comprising detecting in a biological sample from the subject a protein of Pseudomonas aeruginosa, a modified form of the protein or its immunogenic fragment or epitope or an antibody that binds to the protein, where the presence of the protein indicates the infection or exacerbation, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for:

(1) determining the response of a subject suffering from Pseudomonas aeruginosa infection to treatment with a therapeutic compound;  
 (2) diagnosing an acute pulmonary exacerbation in a subject suffering from cystic fibrosis or determining a cystic

fibrosis subject at risk of developing an acute pulmonary exacerbation which comprises diagnosing an infection caused by *Pseudomonas aeruginosa* in the subject, where diagnosis of the infection indicates that the subject is suffering from an acute pulmonary exacerbation or is at risk of developing an acute pulmonary exacerbation;

(3) determining the response of a subject having **cystic fibrosis** and suffering from an acute pulmonary exacerbation to treatment with a therapeutic compound for the exacerbation;

(4) treating a *Pseudomonas aeruginosa* infection in a subject or an acute pulmonary exacerbation in a subject suffering from **cystic fibrosis**;

(5) eliciting the production of an antibody against *Pseudomonas aeruginosa* which comprises administering the protein of *Pseudomonas aeruginosa*;

(6) a vaccine comprising the protein of *Pseudomonas aeruginosa* and a diluent; and

(7) a kit for detecting *Pseudomonas aeruginosa* infection in a biological sample.

ACTIVITY - Antibacterial.

No biological data given.

MECHANISM OF ACTION - Vaccine.

USE - The ferric iron-binding protein (HitA), **thioredoxin** dependent reductase (PAPS), **thioredoxin**, heat shock protein GroES, nucleotide dependent kinase (NDK) or DNA-binding protein HU is useful in the manufacture of a medicament for diagnosing *Pseudomonas aeruginosa* infection or an acute clinical exacerbation. The protein of *Pseudomonas aeruginosa* is useful in preparing a composition for treating or preventing *Pseudomonas aeruginosa* infection. (All claimed.)

Dwg.0/4

FILE SEGMENT: CPI EPI

FIELD AVAILABILITY: AB

MANUAL CODES: CPI: B04-B04B1; B04-B04D4; B04-B04D5; B04-B04G; B04-B04L; B04-F10A6; B04-G07; B04-G21; B04-G22; B04-N03C; B11-C07A; B12-K04A; B12-K04A4B; B14-A01A6; B14-S11B1; B14-S11D3; D05-H04; D05-H07; D05-H09; D05-H11

EPI: S03-E09F; S03-E14H

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ACCESSION NUMBER: 2006-067567 [07] WPIX

DOC. NO. NON-CPI: N2006-058557

DOC. NO. CPI: C2006-024879

TITLE: Detection and/or dosing procedure for anti-transglutaminase antibodies in saliva sample uses immune reaction in pre-treated sample in conditions suitable for formation of immuno-complexes.

DERWENT CLASS: B04 D16 S03

INVENTOR(S): MASCART, F; OCMANT, A

PATENT ASSIGNEE(S): (ULBR) UNIV LIBRE BRUXELLES

COUNTRY COUNT: 111

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA	PG	MAIN	IPC
WO 2005124344	A2 20051229 (200607)*	FR	29	G01N033-53		
RW: AT BE BG BW CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IS IT KE LS LT LU MC MW MZ NA NL OA PL PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW						
W: AE AG AL AM AT AU AZ BA BB BG BR BW BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE EG ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG						

KM KP KR KZ LC LK LS LT LU LV MA MD MG MK MN MW MX MZ NA NG NI  
 NO NZ OM PG PH PL PT RO RU SC SD SE SG SK SL SM SY TJ TM TN TR TT  
 TZ UA UG US UZ VC VN YU ZA ZM ZW

## APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 2005124344	A2	WO 2005-EP6088	20050606

PRIORITY APPLN. INFO: WO 2004-EP6174 20040608

INT. PATENT CLASSIF.:

MAIN: G01N033-53

## BASIC ABSTRACT:

WO2005124344 A UPAB: 20060130

NOVELTY - The procedure for detecting/dosing anti-transglutaminase antibodies in a saliva sample consists of pre-treating the sample with a **mucolytic** compound and then detecting the antibodies by an immune reaction with transglutaminase in conditions that are suitable for the formation of immuno-complexes with the antibodies.

DETAILED DESCRIPTION - The procedure for detecting/dosing anti-transglutaminase antibodies in a saliva sample consists of pre-treating the sample with a **mucolytic** compound and then detecting the antibodies by an immune reaction with transglutaminase in conditions that are suitable for the formation of immuno-complexes with the antibodies. The sampler is one with an activated indicator to show that quantity of the collected sample is adequate, and is selected from the group comprising: Omni-SAL (RTM), Salivette (RTM), Orapette (RTM) and OraSure (RTM). The mucalytic compound is selected from the group comprising: N-acetyl-cystein, nacystelyn, dithiothreitol, gelsolin, **thioredoxin** and EDTA.

USE - Detection of anti-transglutaminase antibodies in saliva for the detection of gluten-induced illnesses such as coeliac disease, or for monitoring a gluten-free regime.

ADVANTAGE - The procedure provides a simple, efficient and non-invasive solution for the diagnosis and monitoring of coeliac disease.

DESCRIPTION OF DRAWING(S) - The drawing shows a diagrammatic representation of the effects of diluting saliva with a preparation containing antigens. (Drawing contains non-English language text)

Dwg.1/11

FILE SEGMENT: CPI EPI

FIELD AVAILABILITY: AB; GI; DCN

MANUAL CODES: CPI: B04-G03; B10-B01B; B10-B02J; B10-E03; B11-C07A;  
 B12-K04A; D05-H09

EPI: S03-E09F; S03-E14H2

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 DUPLICATE 1

ACCESSION NUMBER: 2005:492284 BIOSIS

DOCUMENT NUMBER: PREV200510285061

TITLE: An immunoproteomic approach for identification of clinical biomarkers for monitoring disease - Application to **cystic fibrosis**.

AUTHOR(S): Pedersen, Susanne K. [Reprint Author]; Sloane, Andrew J.; Prasad, Sindhu S.; Sebastian, Lucille T.; Lindner, Robyn A.; Hsu, Michael; Robinson, Michael; Bye, Peter T.; Weinberger, Ron P.; Harry, Jenny L.

CORPORATE SOURCE: Proteome Syst Ltd, 1-35-41 Waterloo Rd, N Ryde, NSW 2113, Australia

SOURCE: sanne.pedersen@proteome-systems.com  
 Molecular & Cellular Proteomics, (AUG 2005) Vol. 4, No. 8,  
 pp. 1052-1060.  
 ISSN: 1535-9476.

DOCUMENT TYPE: Article  
 LANGUAGE: English  
 ENTRY DATE: Entered STN: 16 Nov 2005  
 Last Updated on STN: 16 Nov 2005

ABSTRACT: Circulating antibodies can be used to probe protein arrays of body fluids, prepared by two-dimensional gel electrophoresis, for antigenic biomarker detection. However, detected proteins, particularly low abundance antigens, often remain unidentifiable due to proteome complexity and limiting sample amounts. Using a novel enrichment approach exploiting patient antibodies for isolation of antigenic biomarkers, we demonstrate how immunoproteomic strategies can accelerate biomarker discovery. Application of this approach as a means of identifying biomarkers was demonstrated for \*\*\*cystic\*\*\* fibrosis (CF) lung disease by isolation and identification of inflammatory-associated autoantigens, including myeloperoxidase and calgranulin B from sputum of subjects with CF.

The approach was also exploited for isolation of proteins expressed by the *Pseudomonas aeruginosa* strain PA01. Capture of PA01 antigens using circulating antibodies from CF subjects implicated in vivo expression of *Pseudomonas* proteins. All CF subjects screened, but not controls, were immunoreactive against immunocaptured *Pseudomonas* proteins, representing stress (GroES and ferric iron-binding protein HitA), immunosuppressive (thioredoxin), and alginate synthetase pathway (nucleoside-diphosphate kinase) proteins, implicating their clinical relevance as biomarkers of infection.

CONCEPT CODE: Genetics - Human 03508  
 Clinical biochemistry - General methods and applications 10006  
 Enzymes - General and comparative studies: coenzymes 10802  
 Pathology - Diagnostic 12504  
 Metabolism - Metabolic disorders 13020  
 Digestive system - Pathology 14006  
 Respiratory system - Physiology and biochemistry 16004  
 Respiratory system - Pathology 16006  
 Physiology and biochemistry of bacteria 31000  
 Immunology - General and methods 34502  
 Immunology - Immunopathology, tissue immunology 34508  
 Medical and clinical microbiology - Bacteriology 36002

INDEX TERMS: Major Concepts  
 Infection; Methods and Techniques; Clinical Chemistry (Allied Medical Sciences); Clinical Immunology (Human Medicine, Medical Sciences)

INDEX TERMS: Parts, Structures, & Systems of Organisms  
 sputum: respiratory system

INDEX TERMS: Diseases  
*Pseudomonas aeruginosa* infection: bacterial disease, diagnosis

INDEX TERMS: Diseases  
 cystic fibrosis: respiratory system  
 disease, genetic disease, metabolic disease, digestive system disease, diagnosis  
 Cystic Fibrosis (MeSH)

INDEX TERMS: Chemicals & Biochemicals  
 antibodies; myeloperoxidase [EC 1.11.1.7]; calgranulin B; biomarkers: identification

INDEX TERMS: Methods & Equipment  
 two-dimensional gel electrophoresis: electrophoretic

techniques, laboratory techniques; immunoproteomics:  
laboratory techniques, immunologic techniques

ORGANISM:  
Classifier  
Hominidae 86215  
Super Taxa  
Primates; Mammalia; Vertebrata; Chordata; Animalia  
Organism Name  
human (common): host  
Taxa Notes  
Animals, Chordates, Humans, Mammals, Primates,  
Vertebrates

ORGANISM:  
Classifier  
Pseudomonadaceae 06508  
Super Taxa  
Gram-Negative Aerobic Rods and Cocci; Eubacteria;  
Bacteria; Microorganisms  
Organism Name  
Pseudomonas aeruginosa (species): pathogen, strain-PA01  
Taxa Notes  
Bacteria, Eubacteria, Microorganisms

REGISTRY NUMBER:  
9003-99-0 (myeloperoxidase)  
9003-99-0 (EC 1.11.1.7)

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